# • PCT

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISI	HED U	UN	DER THE PATENT COOPERATION TREATY (PCT)
(51) International Patent Classification 7:		(1	1) International Publication Number: WO 00/23111
A61K 39/395, 48/00, C12P 19/34, C12Q 1/68, G01N 33/53, 33/574, 33/546, 33/567	A1	(4	13) International Publication Date: 27 April 2000 (27.04.00)
(21) International Application Number: PCT/US (22) International Filing Date: 19 October 1999 (			(81) Designated States: CA, JP, US, European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).
(30) Priority Data: 60/104,737 19 October 1998 (19.10.98)  (71) Applicant (for all designated States except US): DIA LLC [US/US]; 3303 Octavius Drive, Santa Clara, (US).	ADEXI		Published  With international search report.  Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.
(72) Inventors; and (75) Inventors/Applicants (for US only): SALCEDA [AR/US]; 4118 Crescendo Avenue, San Jose, C (US). RECIPON, Herve [FR/US]; 85 Fortuna Ave Francisco, CA 94115 (US). CAFFERKEY, Robert Apartment #218, 350 Elan Village Lane, San 3 95134 (US).	CA 951 enue, S t [IE/US	36 San S];	
(74) Agents: LICATA, Jane, Massey et al.; Law Office Massey Licata, 66 E. Main Street, Marlton, NJ 080			
(54) Tide. METUOD OF DIACNOSING MONITODINA	C CTA	CI	NG, IMAGING AND TREATING PROSTATE CANCER
(57) Abstract	U, 31A	101	NG, IMAGING AND TREATING PROSTATE CANCER
	etecting	g, d	iagnosing, monitoring, staging, prognosticating, imaging and treating
			·

# FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
ВВ	Barbados	GH	Ghana	MG	Madagascar	T.J	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	•
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Trinidad and Tobago
BR	Brazil	IL	Israel	MR	Mauritania	UG	Ukraine
BY	Belarus	IS	Iceland	MW	Malawi		Uganda
CA .	Canada	IT	Italy	MX	Mexico	US	United States of America
CF	Central African Republic	JP	Japan	NE	Niger	UZ	Uzbekistan
CG	Congo	KE	Kenya	NL	Netherlands	VN	Viet Nam
СН	Switzerland	KG	Kyrgyzstan	NO		YU	Yugoslavia
CI	Côte d'Ivoire	KP	Democratic People's	NZ	Norway	ZW	Zimbabwe
CM	Cameroon	***	Republic of Korea	PL	New Zealand		
CN	China	KR	Republic of Korea	PT	Poland		
CU	Cuba	KZ	Kazakstan	RO	Portugal		
cz	Czech Republic	LC	Saint Lucia	RU	Romania		
DE	Germany	Ц	Liechtenstein	SD	Russian Federation		
DK	Denmark	LK	Sri Lanka	-	Sudan		
EE	Estonia	LR	Liberia	SE	Sweden		
		LK	Liucia	SG	Singapore		
ľ							

### METHOD OF DIAGNOSING,

### MONITORING, STAGING, IMAGING AND TREATING PROSTATE CANCER

### FIELD OF THE INVENTION

This invention relates, in part, to newly developed 5 assays for detecting, diagnosing, monitoring, staging, prognosticating, imaging and treating cancers, particularly prostate cancer.

### BACKGROUND OF THE INVENTION

Cancer of the prostate is the most prevalent malignancy in adult males, excluding skin cancer, and is an increasingly prevalent health problem in the United States. In 1996, it was estimated that 41,400 deaths would result from this disease in the United States alone, indicating that prostate cancer is second only to lung cancer as the most common cause of death in the same population. If diagnosed and treated early, when the cancer is still confined to the prostate, the chances of cure is significantly higher.

Treatment decisions for an individual are linked to the stage of prostate cancer present in that individual. A common classification of the spread of prostate cancer was developed by the American Urological Association (AUA). The AUA system divides prostate tumors into four stages, A to D. Stage A, microscopic cancer within prostate, is further subdivided into stages A1 and A2. Sub-stage A1 is a well-differentiated cancer confined to one site within the prostate. Treatment is generally observation, radical prostatectomy, or radiation. Sub-stage A2 is a moderately to poorly differentiated cancer at multiple sites within the prostate. Treatment is radical prostatectomy or radiation. Stage B, palpable lump within the prostate, is also further subdivided into sub-stages B1 and B2. In sub-stage B1, the cancer forms a small nodule in one

WO 00/23111

lobe of the prostate. In sub-stage B2, the cancer forms large or multiple nodules, or occurs in both lobes of the prostate. Treatment for sub-stages B1 and B2 is either radical prostatectomy or radiation. Stage C is a large cancer mass 5 involving most or all of the prostate and is also further subdivided into two sub-stages. In sub-stage C1, the cancer forms a continuous mass that may have extended beyond the prostate. In sub-stage C2, the cancer forms a continuous mass that invades the surrounding tissue. Treatment for both these  $10^{\circ}$  sub-stages is radiation with or without drugs to address the The fourth stage, Stage D is metastatic cancer and is also subdivided into two sub-stages. In sub-stage D1, the cancer appears in the lymph nodes of the pelvis. In sub-stage D2, the cancer involves tissues beyond lymph nodes. Treatment 15 for both of these sub-stages is systemic drugs to address the cancer as well as pain.

However, current prostate cancer staging methods are limited. As many as 50% of prostate cancers initially staged as A2, B, or C are actually stage D, metastatic. Discovery of metastasis is significant because patients with metastatic cancers have a poorer prognosis and require significantly different therapy than those with localized cancers. The five year survival rates for patients with localized and metastatic prostate cancers are 93% and 29%, respectively.

Accordingly, there is a great need for more sensitive and accurate methods for the staging of a cancer in a human to determine whether or not such cancer has metastasized and for monitoring the progress of a cancer in a human which has not metastasized for the onset of metastasis.

It has now been found that a number of proteins in the public domain are useful as diagnostic markers for prostate cancer. These diagnostic markers are referred to herein as cancer specific genes or CSGs and include, but are not limited to: Prol09 which is a human zinc-α 2-glycoprotein (Freje et al. Genomics 1993 18(3):575-587); Prol12 which is a human

cysteine-rich protein with a zinc-finger motif (Liebhaber et al. Nucleic Acid Research 1990 18(13):3871-3879; MRO9514772 and WO9845436); Proll1 which is а prostate-specific transglutaminase (Dubbink et al. Genomics 1998 51(3):434-444); 5 Prol15 which is a novel serine protease with transmembrane, LDLR, and SRCR domains and maps to 21q22.3 (Paoloni-Giacobino et al. Genomics 1997 44(3):309-320; WO9837418 and WO987093); Prol10 which is a human breast carcinoma fatty acid synthase (U.S. Patent 5,665,874 and WO9403599); Proll3 which is a 10 homeobox gene, HOXB13 (Steinicki et al. J. Invest. Dermatol. 1998 111:57-63); Prol14 which is a human tetraspan NET-1 (WO9839446); and Prol18 which is a human JM27 protein (WO9845435). ESTs for these CSGs are set forth in SEQ ID NO: 1, 3, 5, 7, 9, 11, 13 and 15 while the full length contigs for 15 these CSGs are set forth in SEQ ID NO:2, 4, 6, 8, 10, 12, 14 and 16, respectively. Additional CSGs for use in the present invention are depicted herein in SEQ ID NO: 17, 18, 19 and 20.

In the present invention, methods are provided for detecting, diagnosing, monitoring, staging, prognosticating, 20 imaging and treating prostate cancer via the cancer specific genes referred to herein as CSGs. For purposes of the present invention, CSG refers, among other things, to native protein expressed by the gene comprising a polynucleotide sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 25 16, 17, 18, 19 or 20. By "CSG" it is also meant herein polynucleotides which, due to degeneracy in genetic coding, comprise variations in nucleotide sequence as compared to SEQ ID NO: 1-20, but which still encode the same protein. alternative, what is meant by CSG as used herein, means the .30 native mRNA encoded by the gene comprising the polynucleotide sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20, levels of the gene comprising the polynucleotide sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 35 20, or levels of a polynucleotide which is capable of

- 4 -

hybridizing under stringent conditions to the antisense sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9,  $\clubsuit$ 0, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20.

Other objects, features, advantages and aspects of the present invention will become apparent to those of skill in the art from the following description. It should be understood, however, that the following description and the specific examples, while indicating preferred embodiments of the invention are given by way of illustration only. Various changes and modifications within the spirit and scope of the disclosed invention will become readily apparent to those skilled in the art from reading the following description and from reading the other parts of the present disclosure.

### SUMMARY OF THE INVENTION

Toward these ends, and others, it is an object of the present invention to provide a method for diagnosing the presence of prostate cancer by analyzing for changes in levels of CSG in cells, tissues or bodily fluids compared with levels of CSG in preferably the same cells, tissues, or bodily fluid type of a normal human control, wherein a change in levels of CSG in the patient versus the normal human control is associated with prostate cancer.

Further provided is a method of diagnosing metastatic prostate cancer in a patient having prostate cancer which is not known to have metastasized by identifying a human patient suspected of having prostate cancer that has metastasized; analyzing a sample of cells, tissues, or bodily fluid from such patient for CSG; comparing the CSG levels in such cells, tissues, or bodily fluid with levels of CSG in preferably the same cells, tissues, or bodily fluid type of a normal human control, wherein an increase in CSG levels in the patient versus the normal human control is associated with prostate cancer which has metastasized.

- 5 -

Also provided by the invention is a method of staging prostate cancer in a human which has such \*cancer by identifying a human patient having such cancer; analyzing a sample of cells, tissues, or bodily fluid from such patient for CSG; comparing CSG levels in such cells, tissues, or bodily fluid with levels of CSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein an increase in CSG levels in the patient versus the normal human control is associated with a cancer which is progressing and a decrease in the levels of CSG is associated with a cancer which is regressing or in remission.

Further provided is a method of monitoring prostate cancer in a human having such cancer for the onset of metastasis. The method comprises identifying a human patient having such cancer that is not known to have metastasized; periodically analyzing a sample of cells, tissues, or bodily fluid from such patient for CSG; comparing the CSG levels in such cells, tissue, or bodily fluid with levels of CSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein an increase in CSG levels in the patient versus the normal human control is associated with a cancer which has metastasized.

Further provided is a method of monitoring the change in stage of prostate cancer in a human having such cancer by looking at levels of CSG in a human having such cancer. The method comprises identifying a human patient having such cancer; periodically analyzing a sample of cells, tissues, or bodily fluid from such patient for CSG; comparing the CSG levels in such cells, tissue, or bodily fluid with levels of CSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein an increase in CSG levels in the patient versus the normal human control is associated with a cancer which is progressing and a decrease in the levels of CSG is associated with a cancer which is regressing or in remission.

Further provided are methods of designing therapeutic agents targeted to a CSG for use in amaging and treating prostate cancer. For example, in one embodiment, therapeutic agents such as antibodies targeted against CSG or 5 fragments of such antibodies can be used to detect or image localization of CSG in a patient for the purpose of detecting or diagnosing a disease or condition. Such antibodies can be polyclonal, monoclonal, or omniclonal or prepared by molecular biology techniques. The term "antibody", as used herein and 10 throughout the instant specification is also meant to include aptamers and single-stranded oligonucleotides such as those derived from an in vitro evolution protocol referred to as SELEX and well known to those skilled in the art. Antibodies can be labeled with a variety of detectable labels including, 15 but not limited to, radioisotopes and paramagnetic metals. Therapeutics agents such as antibodies or fragments thereof can also be used in the treatment of diseases characterized by expression of CSG. In these applications, the antibody can be used without or with derivatization to a cytotoxic agent 20 such as a radioisotope, enzyme, toxin, drug or a prodrug.

Other objects, features, advantages and aspects of the present invention will become apparent to those of skill in the art from the following description. It should be understood, however, that the following description and the specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only. Various changes and modifications within the spirit and scope of the disclosed invention will become readily apparent to those skilled in the art from reading the following description and from reading the other parts of the present disclosure.

# DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to diagnostic assays and methods, both quantitative and qualitative for detecting, diagnosing, monitoring, staging and prognosticating cancers

by comparing levels of CSG in a human patient with those of CSG in a normal human control. For purposes of the present invention, what is meant be CSG levels is, among other things, native protein expressed by the gene comprising a 5 polynucleotide sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20. By "CSG" it is also meant herein polynucleotides which, due to degeneracy in genetic coding, comprise variations in nucleotide sequence as compared to SEQ ID NO: 1-20, but which still encode the 10 same protein. The native protein being detected, may be whole, a breakdown product, a complex of molecules or chemically modified. In the alternative, what is meant by CSG as used herein, means the native mRNA encoded by the gene comprising the polynucleotide sequence of SEQ ID NO:1, 2, 3, 15 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20, levels of the gene comprising the polynucleotide sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20, or levels of a polynucleotide which is capable of hybridizing under stringent conditions to the 20 antisense sequence of SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20. Such levels are preferably determined in at least one of, cells, tissues and/or bodily fluids, including determination of normal and abnormal levels. Thus, for instance, a diagnostic assay in 25 accordance with the invention for diagnosing overexpression of CSG protein compared to normal control bodily fluids, cells, or tissue samples may be used to diagnose the presence of prostate cancer.

All the methods of the present invention may optionally include determining the levels of other cancer markers as well as CSG. Other cancer markers, in addition to CSG, useful in the present invention will depend on the cancer being tested and are known to those of skill in the art.

# Diagnostic Assays

The present invention provides methods for dragnosing the presence of prostate cancer by analyzing for changes in levels of CSG in cells, tissues or bodily fluids compared with levels of CSG in cells, tissues or bodily fluids of preferably the same type from a normal human control, wherein an increase in levels of CSG in the patient versus the normal human control is associated with the presence of prostate cancer.

Without limiting the instant invention, typically, for a quantitative diagnostic assay a positive result indicating the patient being tested has cancer is one in which cells, tissues or bodily fluid levels of the cancer marker, such as CSG, are at least two times higher, and most preferably are at least five times higher, than in preferably the same cells, tissues or bodily fluid of a normal human control.

The present invention also provides a method of diagnosing metastatic prostate cancer in a patient having prostate cancer which has not yet metastasized for the onset of metastasis. In the method of the present invention, a 20 human cancer patient suspected of having prostate cancer which may have metastasized (but which was not previously known to have metastasized) is identified. This is accomplished by a variety of means known to those of skill in the art.

In the present invention, determining the presence of CSG levels in cells, tissues or bodily fluid, is particularly useful for discriminating between prostate cancer which has not metastasized and prostate cancer which has metastasized. Existing techniques have difficulty discriminating between prostate cancer which has metastasized and prostate cancer which has not metastasized and proper treatment selection is often dependent upon such knowledge.

In the present invention, the cancer marker levels measured in such cells, tissues or bodily fluid is CSG, and are compared with levels of CSG in preferably the same cells, tissue or bodily fluid type of a normal human control. That

is, if the cancer marker being observed is just CSG in serum, this level is preferably compared with the lever of CSG in serum of a normal human control. An increase in the CSG in the patient versus the normal human control is associated with prostate cancer which has metastasized.

Without limiting the instant invention, typically, for a quantitative diagnostic assay a positive result indicating the cancer in the patient being tested or monitored has metastasized is one in which cells, tissues or bodily fluid levels of the cancer marker, such as CSG, are at least two times higher, and most preferably are at least five times higher, than in preferably the same cells, tissues or bodily fluid of a normal patient.

Normal human control as used herein includes a human patient without cancer and/or non cancerous samples from the patient; in the methods for diagnosing or monitoring for metastasis, normal human control may preferably also include samples from a human patient that is determined by reliable methods to have prostate cancer which has not metastasized.

### 20 Staging

The invention also provides a method of staging prostate cancer in a human patient. The method comprises identifying a human patient having such cancer and analyzing cells, tissues or bodily fluid from such human patient for CSG. The CSG levels determined in the patient are then compared with levels of CSG in preferably the same cells, tissues or bodily fluid type of a normal human control, wherein an increase in CSG levels in the human patient versus the normal human control is associated with a cancer which is progressing and a decrease in the levels of CSG (but still increased over true normal levels) is associated with a cancer which is regressing or in remission.

### Monitoring

Further provided is a method of monitoring prostate 35 cancer in a human patient having such cancer for the onset of

metastasis. The method comprises identifying a human patient having such cancer that is not known to have metastasized; periodically analyzing cells, tissues or bodily fluid from such human patient for CSG; and comparing the CSG levels determined in the human patient with levels of CSG in preferably the same cells, tissues or bodily fluid type of a normal human control, wherein an increase in CSG levels in the human patient versus the normal human control is associated with a cancer which has metastasized. In this method, normal human control samples may also include prior patient samples.

Further provided by this invention is a method of monitoring the change in stage of prostate cancer in a human patient having such cancer. The method comprises identifying a human patient having such cancer; periodically analyzing cells, tissues or bodily fluid from such human patient for CSG; and comparing the CSG levels determined in the human patient with levels of CSG in preferably the same cells, tissues or bodily fluid type of a normal human control, wherein an increase in CSG levels in the human patient versus the normal human control is associated with a cancer which is progressing in stage and a decrease in the levels of CSG is associated with a cancer which is regressing in stage or in remission. In this method, normal human control samples may also include prior patient samples.

Monitoring a patient for onset of metastasis is periodic and preferably done on a quarterly basis. However, this may be more or less frequent depending on the cancer, the particular patient, and the stage of the cancer.

### Assay Techniques

Assay techniques that can be used to determine levels of gene expression (including protein levels), such as CSG of the present invention, in a sample derived from a patient are well known to those of skill in the art. Such assay methods include, without limitation, radioimmunoassays, reverse transcriptase PCR (RT-PCR) assays, immunohistochemistry

- 11 -

assays, in situ hybridization assays, competitive-binding assays, Western Blot analyses, ELISA assays affd proteomic approaches: two-dimensional gel electrophoresis (2D electrophoresis) and non-gel based approaches such as mass spectrometry or protein interaction profiling. Among these, ELISAs are frequently preferred to diagnose a gene's expressed protein in biological fluids.

An ELISA assay initially comprises preparing an antibody, if not readily available from a commercial source, specific 10 to CSG, preferably a monoclonal antibody. In addition a reporter antibody generally is prepared which binds specifically to CSG. The reporter antibody is attached to a detectable reagent such as radioactive, fluorescent or enzymatic reagent, for example horseradish peroxidase enzyme or alkaline phosphatase.

To carry out the ELISA, antibody specific to CSG is incubated on a solid support, e.g. a polystyrene dish, that binds the antibody. Any free protein binding sites on the dish are then covered by incubating with a non-specific 20 protein such as bovine serum albumin. Next, the sample to be analyzed is incubated in the dish, during which time CSG binds to the specific antibody attached to the polystyrene dish. Unbound sample is washed out with buffer. A reporter antibody specifically directed to CSG and linked to a detectable 25 reagent such as horseradish peroxidase is placed in the dish resulting in binding of the reporter antibody to any monoclonal antibody bound to CSG. Unattached reporter antibody is then washed out. Reagents for peroxidase activity, including a colorimetric substrate are then added -30 to the dish. Immobilized peroxidase, linked to CSG antibodies, produces a colored reaction product. The amount of color developed in a given time period is proportional to the amount of CSG protein present in the sample. Quantitative results typically are obtained by reference to a standard 35 curve.

A competition assay can also be employed wherein antibodies specific to CSG are attached to a solid support and labeled CSG and a sample derived from the host are passed over the solid support. The amount of label detected which is attached to the solid support can be correlated to a quantity of CSG in the sample.

Nucleic acid methods can also be used to detect CSG mRNA as a marker for prostate cancer. Polymerase chain reaction (PCR) and other nucleic acid methods, such as ligase chain 10 reaction (LCR) and nucleic acid sequence based amplification (NASABA), can be used to detect malignant cells for diagnosis and monitoring of various malignancies. For example, reversetranscriptase PCR (RT-PCR) is a powerful technique which can be used to detect the presence of a specific mRNA population 15 in a complex mixture of thousands of other mRNA species. RT-PCR, an mRNA species is first reverse transcribed to complementary DNA (cDNA) with use of the enzyme reverse transcriptase; the cDNA is then amplified as in a standard PCR RT-PCR can thus reveal by amplification the reaction. 20 presence of a single species of mRNA. Accordingly, if the mRNA is highly specific for the cell that produces it, RT-PCR can be used to identify the presence of a specific type of cell.

Hybridization to clones or oligonucleotides arrayed on a solid support (i.e. gridding) can be used to both detect the expression of and quantitate the level of expression of that gene. In this approach, a cDNA encoding the CSG gene is fixed to a substrate. The substrate may be of any suitable type including but not limited to glass, nitrocellulose, nylon or plastic. At least a portion of the DNA encoding the CSG gene is attached to the substrate and then incubated with the analyte, which may be RNA or a complementary DNA (cDNA) copy of the RNA, isolated from the tissue of interest. Hybridization between the substrate bound DNA and the analyte can be detected and quantitated by several means including but

- 13 -

not limited to radioactive labeling or fluorescence labeling of the analyte or a secondary molecule designed to detect the hybrid. Quantitation of the level of gene expression can be done by comparison of the intensity of the signal from the analyte compared with that determined from known standards. The standards can be obtained by in vitro transcription of the target gene, quantitating the yield, and then using that material to generate a standard curve.

Of the proteomic approaches, 2D electrophoresis is a 10 technique well known to those in the art. Isolation of individual proteins from a sample such as accomplished using sequential separation of proteins by different characteristics usually on polyacrylamide gels. First, proteins are separated by size using an electric The current acts uniformly on all proteins, so smaller proteins move farther on the gel than larger proteins. The second dimension applies a current perpendicular to the first and separates proteins not on the basis of size but on the specific electric charge carried by each protein. 20 no two proteins with different sequences are identical on the basis of both size and charge, the result of a 2D separation is a square gel in which each protein occupies a unique spot. Analysis of the spots with chemical or antibody probes, or subsequent protein microsequencing can reveal the relative 25 abundance of a given protein and the identity of the proteins in the sample.

The above tests can be carried out on samples derived from a variety of cells, bodily fluids and/or tissue extracts such as homogenates or solubilized tissue obtained from a patient. Tissue extracts are obtained routinely from tissue biopsy and autopsy material. Bodily fluids useful in the present invention include blood, urine, saliva or any other bodily secretion or derivative thereof. By blood it is meant to include whole blood, plasma, serum or any derivative of blood.

## In Vivo Targeting of CSGs

Identification of these CSGs is also use Tul in the rational design of new therapeutics for imaging and treating cancers, and in particular prostate cancer. For example, in 5 one embodiment, antibodies which specifically bind to CSG can be raised and used in vivo in patients suspected of suffering from prostate cancer. Antibodies which specifically bind a CSG can be injected into a patient suspected of having prostate cancer for diagnostic and/or therapeutic purposes. 10 The preparation and use of antibodies for in vivo diagnosis is well known in the art. For example, antibody-chelators labeled with Indium-111 have been described for use in the radioimmunoscintographic imaging of carcinoembryonic antigen expressing tumors (Sumerdon et al. Nucl. Med. Biol. 1990 15 17:247-254). In particular, these antibody-chelators have been used in detecting tumors in patients suspected of having recurrent colorectal cancer (Griffin et al. J. Clin. Onc. 1991 9:631-640). Antibodies with paramagnetic ions as labels for use in magnetic resonance imaging have also been described 20 (Lauffer, R.B. Magnetic Resonance in Medicine 1991 22:339-Antibodies directed against CSG can be used in a similar manner. Labeled antibodies which specifically bind CSG can be injected into patients suspected of having prostate cancer for the purpose of diagnosing or staging of the disease 25 status of the patient. The label used will be selected in accordance with the imaging modality to be used. For example, radioactive labels such as Indium-111, Technetium-99m or Iodine-131 can be used for planar scans or single photon emission computed tomography (SPECT). Positron emitting 30 labels such as Fluorine-19 can be used in positron emission Paramagnetic ions such as Gadlinium (III) or tomography. Manganese (II) can be used in magnetic resonance imaging (MRI). Localization of the label permits determination of the spread of the cancer. The amount of label within an organ or

- 15 -

tissue also allows determination of the presence or absence of cancer in that organ or tissue.

For patients diagnosed with prostate cancer, injection of an antibody which specifically binds CSG can also have a 5 therapeutic benefit. The antibody may exert its therapeutic effect alone. Alternatively, the antibody can be conjugated to a cytotoxic agent such as a drug, toxin or radionuclide to enhance its therapeutic effect. Drug monoclonal antibodies have been described in the art for example by Garnett and 10 Baldwin, Cancer Research 1986 46:2407-2412. The use of toxins conjugated to monoclonal antibodies for the therapy of various cancers has also been described by Pastan et al. Cell 1986 Yttrium-90 labeled monoclonal antibodies have 47:641-648. been described for maximization of dose delivered to the tumor 15 while limiting toxicity to normal tissues (Goodwin and Meares Cancer Supplement 1997 80:2675-2680). Other cytotoxic radionuclides including, but not limited to Copper-67, Iodineand Rhenium-186 can also be used for labeling of antibodies against CSG.

Antibodies which can be used in these in vivo methods 20 include polyclonal, monoclonal and omniclonal antibodies and antibodies prepared via molecular biology techniques. Antibody fragments and aptamers and single-stranded oligonucleotides such as those derived from an in vitro 25 evolution protocol referred to as SELEX and well known to those skilled in the art can also be used.

Small molecules predicted via computer imaging to specifically bind to regions of CSGs can also be designed and synthesized and tested for use in the imaging and treatment of prostate cancer. Further, libraries of molecules can be screened for potential anticancer agents by assessing the ability of the molecule to bind to CSGs identified herein. Molecules identified in the library as being capable of binding to CSG are key candidates for further evaluation for use in the treatment of prostate cancer.

- 16 -

### **EXAMPLES**

The present invention is further described by the following examples. These examples are provided solely to illustrate the invention by reference to specific embodiments.

These exemplifications, while illustrating certain aspects of the invention, do not portray the limitations or circumscribe the scope of the disclosed invention.

All examples outlined here were carried out using standard techniques, which are well known and routine to those of skill in the art, except where otherwise described in detail. Routine molecular biology techniques of the following example can be carried out as described in standard laboratory manuals, such as Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL, 2nd Ed.; Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1989).

### Example 1: Identification of CSGs

Identification of CSGs were carried out by a systematic analysis of data in the LIFESEQ database available from Incyte Pharmaceuticals, Palo Alto, CA, using the data mining Cancer Leads Automatic Search Package (CLASP) developed by diaDexus LLC, Santa Clara, CA.

The CLASP performs the following steps: selection of highly expressed organ specific genes based on the abundance level of the corresponding EST in the targeted organ versus all the other organs; analysis of the expression level of each highly expressed organ specific genes in normal, tumor tissue, disease tissue and tissue libraries associated with tumor or disease; selection of the candidates demonstrating component ESTs were exclusively or more frequently found in tumor libraries. The CLASP allows the identification of highly expressed organ and cancer specific genes. A final manual in depth evaluation is then performed to finalize the CSGs selection.

- 17 -

Clones depicted in the following Table 1 are CSGs useful in diagnosing, monitoring, staging, imaging and treating prostate cancer.

Table 1: CSGs

5	Clone ID	Pro #	SEQ ID NO:
	3424528H1	Pro109	1,2
	578349H1	Pro112	3,4
	1794013Н1	Proll1	5,6
	2189835Н1	Prol15	7,8
10	3277219Н1	Prol10	9,10
	1857415	Proll3	11,12
	1810463H1	Prol14	13,14
	zr65G11	Prol18	15,16
	2626135H1		17
15	zd46d08		18
	1712252Н1		19
	784583Н1		20

Example 2: Relative Quantitation of Gene Expression

Real-Time quantitative PCR with fluorescent Taqman probes is a quantitation detection system utilizing the 5'-3' nuclease activity of Taq DNA polymerase. The method uses an internal fluorescent oligonucleotide probe (Taqman) labeled with a 5' reporter dye and a downstream, 3' quencher dye.

25 During PCR, the 5'-3' nuclease activity of Taq DNA polymerase releases the reporter, whose fluorescence can then be detected by the laser detector of the Model 7700 Sequence Detection System (PE Applied Biosystems, Foster City, CA, USA).

Amplification of an endogenous control is used to standardize the amount of sample RNA added to the reaction and normalize for Reverse Transcriptase (RT) efficiency. Either cyclophilin, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), ATPase, or 18S ribosomal RNA (rRNA) is used as this endogenous

control. To calculate relative quantitation between all the samples studied, the target RNA levels for one sample were used as the basis for comparative results (calibrator). Quantitation relative to the "calibrator" can be obtained using the standard curve method or the comparative method (User Bulletin #2: ABI PRISM 7700 Sequence Detection System).

The tissue distribution and the level of the target gene were evaluated for every sample in normal and cancer tissues. Total RNA was extracted from normal tissues, cancer tissues, and from cancers and the corresponding matched adjacent tissues. Subsequently, first strand cDNA was prepared with reverse transcriptase and the polymerase chain reaction was done using primers and Taqman probes specific to each target gene. The results were analyzed using the ABI PRISM 7700 Sequence Detector. The absolute numbers are relative levels of expression of the target gene in a particular tissue compared to the calibrator tissue.

# Expression of Clone ID 3424528H1 (Pro109):

For the CSG Prol09, real-time quantitative PCR was 20 performed using the following primers:

Forward Primer:

5'- ATCAGAACAAAGAGGCTGTGTC - 3' (SEQ ID NO:21)
Reverse Primer:

5'- ATCTCTAAAGCCCCAACCTTC - 3' (SEQ ID NO:22)

25 The absolute numbers depicted in Table 2 are relative levels of expression of the CSG referred to as Pro109 in 12 normal different tissues. All the values are compared to normal stomach (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

- 19 -

Table 2: Relative Levels of CSG Pro109 Expression in Pooled
Samples

Tissue	NORMAL
Colon	0.02
Endometrium	0.01
Kidney	0.48
Liver	14.83
Ovary	0.08
Pancreas	4.38
Prostate	11.24
Small Intestine	0.42
Spleen	0
Stomach	1
Testis	0.62
Uterus	0.02

5

10

15

The relative levels of expression in Table 2 show that with the exception of liver (14.83), Pro109 mRNA expression is higher (11.24) in prostate compared with all other normal tissues analyzed. Pancreas, with a relative expression level of 4.38, is the only other tissue expressing considerable mRNA for Pro109.

The absolute numbers in Table 2 were obtained analyzing pools of samples of a particular tissue from different individuals. They cannot be compared to the absolute numbers originated from RNA obtained from tissue samples of a single individual in Table 3.

The absolute numbers depicted in Table 3 are relative levels of expression of Prol09 in 28 pairs of matching samples and 4 unmatched samples. All the values are compared to normal stomach (calibrator). A matching pair is formed by mRNA from the cancer sample for a particular tissue and mRNA from the normal adjacent sample for that same tissue from the same individual.

- 20 -

Table 3: Relative Levels of CSG Pro109 Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matching Normal Adjacent
	Pro34B	Prostate 1	5.98	6.06
5	Pro65XB	Prostate 2	16.68	3.85
	Pro69XB	Prostate 3	20.46	6.82
	Pro78XB	Prostate 4	1.39	1.4
	Pro101XB	Prostate 5	24.8	9.8
	Pro12B	Prostate 6	9.1	0.2
10	Pro13XB	Prostate 7	0.5	9.7
	Pro20XB	Prostate 8	13	12.5
	Pro23B	Prostate 9	16.8	3
	Ovr100050	Ovary 1	0.4	
	Ovr1028	Ovary 2	1.9	
15	Ovr18GA	Ovary 3		0.1
	Ovr206I	Ovary 4		0.1
	Mam12X	Mammary Gland 1	13.5	1.4
	Mam47XP	Mammary Gland 2	0.7	0.2
	Lng47XQ	Lung 1	2.36	0.03
20	Lng60XL	Lung 2	7.39	0.2
	Lng75XC	Lung 3	0.77	0.27
	StoAC44	Stomach 1	0.05	1.19
	StoAC93	Stomach 2	0.55	0.8
	StoAC99	Stomach 3	0.12	3.04
25	ColAS43	Colon 1	16.11	0.07
	ColAS45	Colon 2	0.11	0.08
	ColAS46	Colon 3	4.99	0.4
	Liv15XA	Liver 1	8.43	10.97
	Liv42X	Liver 2	1.57	20.82

- 21 -

Liv94XA	Liver 3	2.98	9.19
Pan77X	Pancreas 1	36	32
Pan82XP	Pancreas 2	0.09	7.09
Pan92X	Pancreas 3	0.7	0
Pan71XL	Pancreas 4	2.48	0.73
Pan10343	Pancreas 5	46	5.5

0 = Negative

5

In the analysis of matching samples, the higher levels of expression were in prostate, showing a high degree of tissue specificity for prostate tissue. Of all the samples different than prostate analyzed, only 4 cancer samples (the cancer sample mammary 1 with 13.5, colon 1 with 16.11, liver 1 with 8.43, and lung 2 with 7.39) showed an expression comparable to the mRNA expression in prostate. These results confirmed some degree of tissue specificity as obtained with the panel of normal pooled samples (Table 2).

Furthermore, the level of mRNA expression was compared in cancer samples and the isogenic normal adjacent tissue from the same individual. This comparison provides an indication of specificity for the cancer (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent). Table 3 shows overexpression of Prolo9 in 6 out of 9 primary prostate cancer tissues compared with their respective normal adjacents. Thus, overexpression in the cancer tissue was observed in 66.66% of the prostate matching samples tested (total of 9 prostate matching samples).

Altogether, the degree of tissue specificity, plus the mRNA overexpression in 66.66% of the primary prostate matching samples tested is indicative of Prol09 being a diagnostic 30 marker for prostate cancer.

- 22 -

# Expression of Clone ID 578349H1 (Pro112):

For the CSG Proll2, real-time quantitative PCR was performed using the following primers:

Forward Primer

5'- TGCCGAAGAGGTTCAGTGC - 3' (SEQ ID NO:23)

Reverse Primer

5'- GCCACAGTGGTACTGTCCAGAT - 3' (SEQ ID NO:24)

The absolute numbers depicted in Table 4 are relative levels of expression of the CSG Pro112 in 12 normal different tissues. All the values are compared to normal thymus (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

Table 4: Relative Levels of CSG Prol12 Expression in Pooled

Samples

Tissue	NORMAL
Brain	2.9
Heart	0.1
Kidney	0.2
Liver	0.2
Lung	7.7
Mammary	4.2
Muscle	0.1
Prostate	5.5
Small Intestine	1.8
Testis	1
Thymus	1
Uterus	21

25

20

5

The relative levels of expression in Table 4 show that

30 Proll2 mRNA expression is the 3<sup>rd</sup> most highly expressed gene
(after uterus and mammary) in the pool of normal prostate
tissue compared to a total of 12 tissues analyzed. The
absolute numbers in Table 4 were obtained analyzing pools of
samples of a particular tissue from different individuals.

35 These results demonstrate that Proll2 mRNA expression is specific for prostate thus indicating Proll2 to be a diagnostic marker for prostate disease especially cancer.

- 23 -

### Expression of Clone ID 1794013H1 (Prol11):

For the CSG Prolll, real-time quantitative PCR was performed using the following primers:

Forward Primer

5'- GCTGCAAGTTCTCCACATTGA - 3' (SEQ ID NO:25)

Reverse Primer

5'- CAGCCGCAGGTGAAACAC - 3' (SEQ ID NO:26)

The absolute numbers depicted in Table 5 are relative levels of expression of the CSG Proll1 in 12 normal different tissues. All the values are compared to normal testis (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

Table 5: Relative Levels of CSG Proll1 Expression in Pooled Samples

Tissue	NORMAL
Brain	0.04
Heart	0
Kidney	0
Liver	0
Lung	0.05
Mammary	0.14
Muscle	5166.6
Prostate	1483.72
Small Intestine	0.33
Testis	1
Thymus	0.49
Uterus	0.07

25

20

15

5

The relative levels of expression in Table 5 show that Proll1 mRNA expression is extraordinarily high in the pool of normal prostate (1483.72) compared to all the other tissues analyzed with the exception of muscle (5166.6). These results demonstrate that Proll1 mRNA expression shows specificity for prostate and muscle.

The absolute numbers in Table 5 were obtained analyzing pools of samples of a particular tissue from different

individuals. They cannot be compared to the absolute numbers originated from RNA obtained from tissue samples of a single individual in Table 6.

The absolute numbers depicted in Table 6 are relative 5 levels of expression of Proll1 in 48 pairs of matching and 18 unmatched samples. All the values are compared to normal testis (calibrator). A matching pair is formed by mRNA from the cancer sample for a particular tissue and mRNA from the normal adjacent sample for that same tissue from the same 10 individual.

Table 6: Relative Levels of CSG Proll1 Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matching Normal Adjacent
	Pro101XB	Prostate 1	8.3	21.8
15	Pro12B	Prostate 2	2336	133
	Pro13XB	Prostate 3	3.4	23
	Pro20XB	Prostate 4	21.6	121.5
	Pro23B	Prostate 5	19.4	3.7
	Pro34B	Prostate 6	15	39
20	Pro65XB	Prostate 7	8	867
	Pro69XB	Prostate 8	56	94
	Pro78XB	Prostate 9	24	1515
	Pro84XB	Prostate 10	119	15.35
	Pro90XB	Prostate 11	8.08	112.2
25	Pro91XB	Prostate 12	0.88	51.8
-	ProC215	Prostate 13	0.3	
	ProC234	Prostate 14	0.35	
	ProC280	Prostate 15	436.5	
	Pro109XB	Prostate 16	3.43	265
30	Proll0	Prostate 17	18.2	8.73

			<del></del>	<del></del>
	Pro125XB	Prostate 18	0.34	186
	Pro326	Prostate 19	1392	110
	Pro10R	Prostate 20 (prostatitis)	0.5	
	Pro20R	Prostate 21 (prostatitis)	24.1	
5	Pro258	Prostate 22 (BPH)	4610	
	Pro263C	Prostate 23 (BPH)	0	
	Pro267A	Prostate 24 (BPH)	1.46	
	Pro271A	Prostate 25 (BPH)	0	
	Pro460Z	Prostate 26 (BPH)	1.47	
10	ProC032	Prostate 27 (BPH)	14.4	
	Tst39X	Testis 1	0	0
	Bld32XK	Bladder 1	0.44	0.41
	Bld46XK	Bladder 2	0	0
	Bld66X	Bladder 3	0	0
15	BldTR14	Bladder 4	0	0
	Kid106XD	Kidney 1	0	0
	Kid107XD	Kidney 2	0	0
	Kid109XD	Kidney 3	0	0
	Pan10343	Pancreas 1	0	0
20	Pan71XL	Pancreas 2	0	0
	Pan77X	Pancreas 3	0	0
	Liv15XA	Liver 1	0	0
	Liv42X	Liver 2	0	0
	ClnAS43	Colon 1	0	0
25	ClnAS45	Colon 2	0	0
	ClnAS46	Colon 3	0	0
	ClnAS67	Colon 4	0 .	0
	ClnAC19	Colon 5	0	0
	ClnAS12	Colon 6	0	0

- 26 -

	SmI21XA	Small Intestine 1	0	0
	SmIH89	Small Intestine 2	0	0
	Lng47XQ	Lung 1	0.7	0
	Lng60XL	Lung 2	0	0
5	Lng75XC	Lung 3	0	0
	Lng90X	Lung 4	0	0
	Mam12X	Mammary Gland 1	0	1.4
	Mam59X	Mammary Gland 2	0.2	0
	MamA06X	Mammary Gland 3	0	0
10	MamS127	Mammary Gland 4	0	0
	Mam162X	Mammary Gland 5	0	0
	Mam42DN	Mammary Gland 6	0	0
	Ovr103X	Ovary 1	0.14	0
	Ovr10050	Ovary 2	0.2	
15	Ovr1028	Ovary 3	0	
	Ovr10400	Ovary 4	0.2	
	Ovr18GA	Ovary 5		0
	Ovr2061	Ovary 6		0
	Ovr20GA	Ovary 7		0.2
20	Ovr25GA	Ovary 8		0
	A 11			L I

0= Negative

In the analysis of matching samples, the higher levels of expression were in prostate showing a high degree of tissue specificity for prostate. These results confirm the tissue specificity results obtained with normal pooled samples (Table 5).

Furthermore, the level of mRNA expression in cancer samples and the isogenic normal adjacent tissue from the same individual were compared. This comparison provides an indication of specificity for cancer (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent). Table 6 shows overexpression of Prolll in 5 out

- 27 -

of 16 primary prostate cancer samples compared with their respective normal adjacent (prostate samples 2, 5, 10, 17, and 19). Similar expression levels were observed in 3 unmatched prostate cancers (prostate samples 13, 14, 15), 2 prostatitis (prostate samples 20, 21), and 6 benign prostatic hyperplasia samples (prostate samples 22 through 27). Thus, there is overexpression in the cancer tissue of 31.25% of the prostate matching samples tested (total of 16 prostate matching samples).

Altogether, the high level of tissue specificity, plus the mRNA overexpression in 31.25% of the prostate matching samples tested are indicative of Proll1 being a diagnostic marker for prostate cancer.

# Expression of Clone ID 2189835H1 (Pro115):

15 For the CSG Proll5, real-time quantitative PCR was performed using the following primers:

Forward Primer

5'- TGGCTTTGAACTCAGGGTCA - 3' (SEQ ID NO:27)

Reverse Primer

20 5'- CGGATGCACCTCGTAGACAG - 3' (SEQ ID NO:28)

The absolute numbers depicted in Table 7 are relative levels of expression of the CSG Prol15 in 12 normal different tissues. All the values are compared to normal thymus (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

Table 7: Relative Levels of CSG Prol15 Expression in Pooled Samples

Tissue	NORMAL
Brain	0.016
Heart	0.002
Kidney	8.08
Liver	2.20
Lung	112.99

30

-	2	R	_

Mammary	29.45
Muscle	0.05
Prostate	337.79
Small Intestine	7.54
Testis	1.48
Thymus	1
Uterus	1.4

5

The relative levels of expression in Table 7 show that Prol15 mRNA expression is higher (337.79) in prostate compared with all the other normal tissues analyzed. Lung, with a relative expression level of 112.99, and mammary (29.446) are the other tissues expressing moderate levels of mRNA for Prol15. These results establish Prol15 mRNA expression to be highly specific for prostate.

The absolute numbers in Table 7 were obtained analyzing pools of samples of a particular tissue from different individuals. They cannot be compared to the absolute numbers originated from RNA obtained from tissue samples of a single individual in Table 8.

The absolute numbers depicted in Table 8 are relative levels of expression of Prol15 in 17 pairs of matching and 21 unmatched samples. All the values are compared to normal thymus (calibrator). A matching pair is formed by mRNA from the cancer sample for a particular tissue and mRNA from the 25 normal adjacent sample for that same tissue from the same individual.

Table 8: Relative Levels of CSG Prol15 Expression in Individual Samples

_	Sample ID	Tissue	Cancer	Matching Normal Adjacent
30	Pro12B	Prostate 1	1475.9	190.3
	ProC234	Prostate 2	169.61	
	Pro109XB	Prostate 3		639.53
	Pro101XB	Prostate 4	1985.2	2882.9

WO 00/23111

- 29 -

	Pro13XB	Prostate 5	34.9	13.9
	Pro215	Prostate 6	525.59	
	Pro125XB	Prostate 7		556.05
	Pro23B	Prostate 8	1891.4	1118.6
5	ProC280	Prostate 9	454.3	
	Pro20XB	Prostate 10	1332.6	
	Pro34B	Prostate 11		362.91
	Pro65XB	Prostate 12		135.06
	Pro69XB	Prostate 13		179.67
10	Pro10R	Prostate 14 (prostatitis)	143.82	
	Pro20R	Prostate 15 (prostatitis)	397.79	
	Pro258	Prostate 16 (BPH)	216.6	
	Pro263C	Prostate 17 (BPH)	601.25	
	Pro267A	Prostate 18 (BPH)	200.28	
15	Pro271A	Prostate 19 (BPH)	111.43	
	Pro460Z	Prostate 20 (BPH)	53.84	
	ProC032	Prostate 21 (BPH)	56.94	
	SmI21XA	Small Intestine 1	28.8	29.9
	SmIH89	Small Intestine 2	70.8	348.5
20	ClnAC19	Colon 1	22.73	446.47
	ClnAS12	Colon 2	116.97	493.18
	Kid106XD	Kidney 1	86.13	41.14
	Kid107XD	Kidney 2	0.26	35.14
	Lng47XQ	Lung 1	5.13	20.98
25	Lng60XL	Lung 2	13.93	114.78
!	Lng75XC	Lung 3	16.47	53.79
	Mam12X	Mammary Gland 1	6.25	10.75
	Mam162X	Mammary Gland 2	1.84	2.54
	Mam42DN	Mammary Gland 3	23.08	35.51

Ovr10050	Ovary 1	0.9	
Ovr1028	Ovary 2	261.4	*
Ovr103X	Ovary 3	7	0.1
Ovr20GA	Ovary 4		0
Ovr25GA	Ovary 5		0

0 = Negative

5

Higher levels of expression were seen in prostate, showing a high degree of tissue specificity for prostate 10 tissue. Of all the analyzed samples different from prostate, only two cancer samples (colon 2 with 116.97 and ovary 2 with 261.4), and 5 normal adjacent tissue samples (small intestine 2, colon 1, colon 2, kidney 1, and lung 2), showed an expression comparable to the mRNA expression in prostate. 15 These results confirmed the tissue specificity results

obtained with the panel of normal pooled samples (Table 7).

Furthermore, the levels of mRNA expression in cancer samples and the isogenic normal adjacent tissue from the same individual were compared. This comparison provides an indication of specificity for the cancer (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent). Table 8 shows higher expression of Prol15 in 3 out of 4 matched prostate cancer tissues (prostate samples 1, 5 & 8).

Altogether, the high level of tissue specificity, plus the higher expression in 75% of the prostate matching samples tested, are indicative of Prol15 being a diagnostic marker for prostate cancer.

# Expression of Clone ID 3277219H1 (Prol10):

30 For the CSG ProllO, real-time quantitative PCR was performed using the following primers:

Forward Primer

5'- CGGCAACCTGGTAGTGAGTG - 3' (SEQ ID NO:29)

- 31 -

### Reverse Primer

5'- CGCAGCTCCTTGTAAACTTCAG - 3' (SEQ 19 NO:30)

The absolute numbers depicted in Table 9 are relative levels of expression of the CSG Prol10 in 12 normal different 5 tissues. All the values are compared to normal small intestine (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

Table 9: Relative Levels of CSG Prol10 Expression in Pooled
Samples

Tissue	NORMAL
Brain	6.61
Heart	0.7
Kidney	0.74
Liver	7.94
Lung	11.88
Mammary	22.78
Muscle	6.77
Prostate	3.01
Small Intestine	1
Testis	2.58
Thymus	13.74
Uterus	2.61

The relative levels of expression in Table 9 show that Prol10 25 mRNA expression is not as high in normal prostate (3.01) compared with all the other normal tissues analyzed.

The absolute numbers in Table 9 were obtained analyzing pools of samples of a particular tissue from different individuals. They cannot be compared to the absolute numbers originated from RNA obtained from tissue samples of a single individual in Table 10.

The absolute numbers depicted in Table 10 are relative levels of expression of Prol10 in 33 pairs of matching samples. All the values are compared to normal small intestine (calibrator). A matching pair is formed by mRNA from the cancer sample for a particular tissue and mRNA from

15

10

20

WO 00/23111

- 32 -

the normal adjacent sample for that same tissue from the same individual.

Table 10: Relative Levels of CSG Prol10 Expression in Individual Samples

5	Sample ID	Tissue	Cancer	Matching Normal Adjacent
	Pro12B	Prostate 1	11.8	0.3
	Pro78XB	Prostate 2	14.3	6.3
	Pro101XB	Prostate 3	33.2	10.7
	Pro13XB	Prostate 4	0.3	0.4
10	Pro23XB	Prostate 5	25.5	14.4
	Pro20XB	Prostate 6	43.3	4
	Pro34XB	Prostate 7	31.8	18.7
	Pro65XB	Prostate 8	26.9	3.4
	Pro69XB	Prostate 9	12.5	7
15	Lng75XC	Lung 1	1.9	3
	Lng90X	Lung 2	5.5	0.5
	LngAC11	Lung 3	9.3	9.7
	LngAC32	Lung 4	11.2	2.2
	Lng47XQ	Lung 5	11.3	0.3
20	Lng60XL	Lung 6	29.1	6.8
	Mam12B	Mammary Gland 1	19.8	0
	Mam603X	Mammary Gland 2	13.7	0
	Mam82XI	Mammary Gland 3	73.5	0
	MamA04	Mammary Gland 4	0	24.6
25	MamB011X	Mammary Gland 5	17.4	2
	MamC012	Mammary Gland 6	0	12.8
	MamC034	Mammary Gland 7	0	61
	Mam12X	Mammary Gland 8	14	2.2
	Mam59X	Mammary Gland 9	33	2.2

- 33 -

MamA06X	Mammary Gland 10	16.4	0.8
Liv15XA	Liver 1	4.7	0.6
Liv42X	Liver 2	7.5	2.6
Liv94XA	Liver 3	0.4	1.4
ClnAS43	Colon 1	52.9	1.4
ClnAS45	Colon 2	2.1	0.8
ClnAS46	Colon 3	39.8	3.7
SmI21X	Small Intestine 1	0.9	0.1
SmIH89	Small Intestine 2	5.8	0.9

10 = Negative

5

The levels of mRNA expression in cancer samples and the isogenic normal adjacent tissue from the same individual were compared. This comparison provides an indication of specificity for the cancer (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent). Table 10 shows overexpression of Prol10 in 8 of the 9 primary prostate cancer tissues compared with their respective normal adjacent (except prostate 4). Thus, there was overexpression in 88.88% of the cancer prostate tissue as compared to the prostate matching samples tested (total of 9 prostate matching samples).

Although not tissue specific, Prol10 mRNA expression is upregulated in prostate cancer tissues. The mRNA overexpression in 88.88% of the primary prostate matching 25 cancer samples tested is indicative of Prol10 being a diagnostic marker for prostate cancer. Prol10 also showed overexpression in several other cancers tested including small intestine, colon, liver, mammary and lung (see Table 10). Accordingly Prol10 may be a diagnostic marker for other types of cancer as well.

- 34 -

# Expression of Clone ID 1857415; Gene ID 346880 (Proll3):

For the CSG Proll3, real-time quantitati  $\P$ e PCR was performed using the following primers:

Forward Primer

5'- CGGGAACCTACCAGCCTATG - 3' (SEQ ID NO:31)

Reverse Primer

5

5'- CAGGCAACAGGGAGTCATGT - 3' (SEQ ID NO:32)

The absolute numbers depicted in Table 11 are relative levels of expression of the CSG Prol13 in 12 normal different tissues. All the values are compared to normal thymus (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

Table 11: Relative Levels of CSG Pro113 Expression in
Pooled Samples

	Tissue	NORMAL
	Brain	0.03
	Heart	0
	Kidney	0.01
20	Liver	0
	Lung	0
	Mammary Gland	0
	Muscle	0.04
	Prostate	489.44
25	Small Intestine	0.02
	Testis	0.35
	Thymus	1
	Uterus	0.13

The relative levels of expression in Table 11 show that Prol13

30 mRNA expression is higher (489.44) in prostate compared with all the other normal tissues analyzed. Testis, with a relative expression level of 0.35, uterus (0.13), thymus (1.0), kidney (0.01) and brain (0.03) were among the other tissues expressing lower mRNA levels for Prol13. These results establish that Prol13 mRNA expression is highly specific for prostate.

- 35 -

The absolute numbers in Table 11 were obtained analyzing pools of samples of a particular tissue from different individuals. They cannot be compared to the absolute numbers originated from RNA obtained from tissue samples of a single 5 individual in Table 12.

The absolute numbers depicted in Table 12 are relative levels of expression of Prol13 in 78 pairs of matching and 25 unmatched tissue samples. All the values are compared to normal thymus (calibrator). A matching pair is formed by mRNA 10 from the cancer sample for a particular tissue and mRNA from the normal adjacent sample for that same tissue from the same individual. In cancers (for example, ovary) where it was not possible to obtain normal adjacent samples from the same individual, samples from a different normal individual were analyzed.

Table 12: Relative Levels of CSG Prol13 Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matched or Unmatched Normal Adjacent
	Pro780B/781B	Prostate 1	375.58	446.29
20	Pro1291B/1292B	Prostate 2	1060	31
	Pro139B96/140B96	Prostate 3	41	32
	Pro209B96/210B96	Prostate 4	505	255
	Pro1256B/1257B	Prostate 5	165.79	141.63
	Pro1293B/1294B	Prostate 6	1613.7	874.61
25	Pro694B/695B	Prostate 7	458.6	142.21
	Pro1012B/1013B	Prostate 8	1520	864
•	Pro1222B/1223B	Prostate 9	939	530
	Pro845B/846B	Prostate 10	1552.4	374.6
	Pro1094B/1095B	Prostate 11	278.37	135.89
30	Pro650B/651B	Prostate 12	532.81	640.85

	Pro902B/903B	Prostate 13	609.05	415.86
	Pro916B/917B	Prostate 14	699.42	401.24
	Pro9821110A/110B	Prostate 15	156	487.8
	ProS9821326A/26B	Prostate 16	744.4	472.8
5	Pro9407c215	Prostate 17	1389.2	
	Pro9407c234	Prostate 18	305.5	
	Pro9407c280A	Prostate 19	894.5	
	Pro9409C010R	Prostate 20 (prostatitis)	269.7	
	Pro9404C120R	Prostate 21 (prostatitis)	299.2	
10	Pro1000258	Prostate 22 (BPH)	149.6	
	Pro4001263C	Prostate 23 (BPH)	576	
	Pro4001267A	Prostate 24 (BPH)	132.1	
	Pro9411C032	Prostate 25 (BPH)	118.2	
	Pro4001460Z	Prostate 26 (BPH)	276.3	
15	Pro4001271A	Prostate 27 (BPH)	58.7	
	Kid1064D/65D	Kidney 1	0	0.1
	Kid1079D/1080D	Kidney 2	0.3	0.02
	Kid1097D/1098D	Kidney 3	35.14	0.32
	Kid1024D/1025D	Kidney 4	1.31	0
20	Kid1183D/1184D	Kidney 5	24.79	0
	Kid1242D/1243D	Kidney 6	0	0
•	Bld469K	Bladder 1		2.88
	Bld467K/468K	Bladder 2	2.65	
	Bld327K/328K	Bladder 3	0	4.05
25	Bld470K	Bladder 4		1.64
	Bld665T/664T	Bladder 5	0.21	1.99

	Bld1496K/1497K	Bladder 6	13.55	1.14
	Bld1721K/1722K	Bladder 7	120.16	1.34
	Tst239X/240X	Testis 1	31.5	0.73
	TstS9820647A/47B	Testis 2	15.7	0
5	TstS9820663A/663B	Testis 3	72	1.4
	SknS9821248A/248B	Skin 1	1.8	0.5
	SknS99448A/448B	Skin 2	251.6	0
	Skn99816A/816B	Skin 3	33	0.7
	Sto4004864A4/B4	Stomach 1	14.12	0
10	Sto4004509A3/B1	Stomach 2	40.74	39
	SmI9807A212A/213A	Small Intestine 1	0.1	0
	SmI9802H008/H009	Small Intestine 2	5.8	0.1
	Cln9608B012/B011	Colon 1	4.5	0
	Cln9709c074ra/073ra	Colon 2	65.8	3.1
15	Cln4004709A1/709B1	Colon 3	1.1	0.9
	Cln9405C199/C200	Colon 4	34.76	0.73
	Cln9707c004gb/006ga	Colon 5	90.26	0.96
	Cln96-09-B004/B003	Colon 6	17.9	20.64
	Cln9612B006/B005	Colon 7	17.56	0.3
20	Cln9705F002D/F001C	Colon 8	21.39	0
	ClnCXGA	Colon 9	429.14	142.69
	Pan10343a	Pancreas 1	0	0
	Pan776P/777P	Pancreas 2	0	0.15
	Pan9210/9220	Pancreas 3	7.36	0
25	Pan714L/715L	Pancreas 4	13.57	0.11
	Pan824P/825P	Pancreas 5	0	0
	Lng476Q/477Q	Lung 1	0	0
	Lng605L/606L	Lung 2	0	0.1
	Lng11145B/11145C	Lung 3	85.9	0

	Lng0008632A/32B	Lung 4	22.05	
	Lng750C/751C	Lung 5	23.85	0
	Lng8890A/8890B	Lung 6	10.63	0.25
	Lng8926A/8926B	Lung 7	<del>                                     </del>	0
5		Lung 8	15.37	0
	Lng9502C109R/110R	Lung 9	26.17	0
	LngS9821944a/44b		0.68	0
	Mam00042D01/42N01	Lung 10	0	0
	Mam59XC	Mammary Gland 1	8.5	0
10	Mam9706A066G/67C	Mammary Gland 2	61.07	0
10	Mam14153a1C	Mammary Gland 3	4.84	0
		Mammary Gland 4	9.72	6.99
	Mam1620F/1621F	Mammary Gland 5	0.91	0
	Mam00014D05	Mammary Gland 6	2.45	0
	End10479B/D	Endometrium 1	133.43	1.12
15	End9705A125A/126A	Endometrium 2	0	0.39
	End9704C281A/282A	Endometrium 3	23.5	1.56
	End680o97/681o97	Endometrium 4	88.89	79.02
	Utr13590/13580	Uterus 1	0.2	0
	Utr850U/851U	Uterus 2	0	0
20	Utr14170/14180	Uterus 3	14	0.4
	Utr233U96/234U96	Uterus 4	8.65	4.64
	CvxVNM00052D01/52N01	Cervix 1	0.82	77.15
	CvxVNM00083D01/83N01	Cervix 2	0.78	221.48
	CvxND00023D01/23N01	Cervix 3	3.25	15.22
25	Ovr10370/10380	Ovary 1	0.1	0
_	Ovr10050	Ovary 2	18.96	
	Ovr1028	Ovary 3	0	
	Ovr14638A1C	Ovary 4	3.2	
	Ovr14603A1D	Ovary 5	882.3	
30	Ovr7730	Ovary 6	0	

_	3	9	_

Ovr9702C018GA	Ovary 7	0.15
Ovr206I	Ovary 8	0
Ovr9702C020GA	Ovary 9	0
Ovr9702C025GA	Ovary 10	0
Ovr9701C035GA	Ovary 11	0.07
Ovr9701C050GB	Ovary 12	0.58

0 = Negative

5

In the analysis of matching samples, the higher levels of expression were in prostate, showing a high degree of 10 tissue specificity for prostate tissue. In addition to the higher expression levels in prostate cancer samples, Proll3 expression was found to be either induced (where not expressed in normal adjacent tissues) or somewhat upregulated in several other cancers. However, the relative expression and the fold increase in prostate cancer samples far exceeds that in other cancer tissues and is highly significant.

Furthermore, the levels of mRNA expression in cancer samples and the isogenic normal adjacent tissue from the same individual were compared. This comparison provides an 20 indication of specificity for the cancer (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent). Table 12 shows overexpression of Prol13 in 13 out of 16 primary prostate cancer tissues compared with their respective normal adjacent (prostate samples 2, 3, 4, 5, 6 7, 25 8, 9, 10, 11, 13, 14, 16). Thus, there was overexpression in the cancer tissue for 81.25% of the prostate matching samples tested. The median for the level of expression in prostate cancer tissue samples is 609, whereas the median for all other cancers is only 7.93, with the exception of one colon sample, 30 colon 9, whose expression was similar to that found in prostate cancer tissues.

Altogether, the high level of tissue specificity, plus the mRNA overexpression in 81.25% of the primary prostate matching samples tested are indicative of Proll3 being a

diagnostic marker for prostate cancer. Expression was also found to be higher in other cancer tissues compared with their respective normal adjacent tissues (kidney, bladder, testis, skin, stomach, small intestine, colon, pancreas, lung, mammary, endometrium, uterus, and ovary) thus indicating Proll3 to be a pan cancer marker.

## Expression of Clone ID 1810463H1 (Pro114):

For the CSG Proll4, real-time quantitative PCR was performed using the following primers:

10 Forward Primer

5'- TGGGCATCTGGGTGTCAA - 3' (SEQ ID NO:33)

Reverse Primer

5'- CGGCTGCGATGAGGAAGTA - 3' (SEQ ID NO:34)

The absolute numbers depicted in Table 13 are relative levels of expression of the CSG Prol14 in 12 normal different tissues. All the values are compared to normal muscle (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

20 Table 13: Relative Levels of CSG Proll4 Expression in Pooled Samples

	Tissue	NORMAL
	Brain	9.7
	Heart	0.7
25	Kidney	414.4
	Liver	4
•	Lung	882.2
	Mammary	44
	Muscle	1
30	Prostate	1951
	Small Intestine	22
	Testis	367.1
	Thymus	25.8
	Uterus	139.6

35 The relative levels of expression in Table 13 show that Prol14 mRNA expression is higher (1951) in prostate compared with all the other normal tissues analyzed. Lung, with a relative

- 41 -

expression level of 882.2, kidney 414.4, testis 367.1 and uterus 139.6, are the other tissues expressing higher levels of mRNA for Proll4. These results establish Proll4 mRNA expression to be more specific for prostate than other tissues 5 examined.

The high level of tissue specificity is indicative of Proll4 being a diagnostic marker for diseases of the prostate, especially cancer.

### Expression of Clone ID zr65g11 (Pro118):

10 For the CSG Prol18, real-time quantitative PCR was performed using the following primers:

Forward Primer

5'- GCCCATCTCCTGCTTCTTTAGT - 3' (SEQ ID NO:35)

Reverse Primer

15 5'- CGTGGAGATGGCTCTGATGTA - 3' (SEQ ID NO:36)

The absolute numbers depicted in Table 14 are relative levels of expression of the CSG Prol18 in 12 normal different tissues. All the values are compared to normal kidney (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

Table 14: Relative Levels of CSG Prol18 Expression in Pooled Samples

	Tissue	NORMAL
25	Colon	0.87
	Endometrium	19282
	Kidney	1
_	Liver	0
	Ovary	86.22
30	Pancreas	0
	Prostate	962.1
	Small Intestine	0
	Spleen	0.75
	Stomach	0.54
35	Testis	343.7
	Uterus	1064

The relative levels of expression in Table \$4 show that Prol18 mRNA expression is the  $3^{rd}$  highest in prostate (962.1) next to endometrium (19282) and uterus (1064), which are female-specific tissues. Testis, with a relative expression 5 level of 343.7 is the only other male tissue expressing moderate levels of mRNA for Prol18. These results establish Prol18 mRNA expression to be highly specific for reproductive tissues including the prostate.

The absolute numbers in Table 14 were obtained analyzing 10 pools of samples of a particular tissue from different individuals. They cannot be compared to the absolute numbers originated from RNA obtained from tissue samples of a single individual in Table 15.

The absolute numbers depicted in Table 15 are relative 15 levels of expression of Prol18 in 59 pairs of matching and 21 unmatched samples. All the values are compared to normal kidney (calibrator). A matching pair is formed by mRNA from the cancer sample for a particular tissue and mRNA from the normal adjacent sample for that same tissue from the same 20 individual.

Table 15: Relative Levels of CSG Prol18 Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matching Normal Adjacent
	Pro12B	Prostate 1	41700.7	22242.83
25	ProC234	Prostate 2	40087	
	Pro78XB	Prostate 3	4075.6	7066.7
-	Pro109XB	Prostate 4	334.4	777.2
	Pro84XB	Prostate 5	11684	58290
	Pro101XB	Prostate 6	21474.13	100720.8
30	Pro91X	Prostate 7	14849	33717
`	Pro13XB	Prostate 8	202.57	146.91

•				
	ProC215	Prostate 9	73243	
	Pro125XB	Prostate 10	629.6	521.4
	Pro23B	Prostate 11	157532.6	110654.4
	Pro90XB	Prostate 12	2317	64134
5	ProC280	Prostate 13	42020	
	Pro20XB	Prostate 14	2909.31	
	Pro34B	Prostate 15	29610	23264
	Proll0	Prostate 16	13354	30991
	Pro65XB	Prostate 17	10126	11270
10	Pro69XB	Prostate 18		2671.42
	Pro326	Prostate 19	9962.3	19231
	Pro10R	Prostate 20 (prostatitis)	27355	
	Pro20R	Prostate 21 (prostatitis)	21081	
	Pro258	Prostate 22 (BPH)	79916.32	
15	Pro263C	Prostate 23 (BPH)	108924.5	_
	Pro267A	Prostate 24 (BPH)	92910.22	
	Pro271A	Prostate 25 (BPH)	57004.4	
	Pro460Z	Prostate 26 (BPH)	57449.23	
	ProC032	Prostate 27 (BPH)	45781.44	
20	Kid106XD	Kidney 1	3.08	217.36
	Kid107XD	Kidney 2	0	38.36
	Kid109XD	Kidney 3	0	123.5
	Kid10XD	Kidney 4	17.69	67.8
	Kid11XD	Kidney 5	16.74	360.8
. 25	Kid124D	Kidney 6	0	167.4
	Bld32XK	Bladder 1	0	0
	Bld47K	Bladder 2		36.38
	Bld66X	Bladder 3	0	4.52
	BldTR14	Bladder 4	0	12.17

BldTR17	Bladder 5	0	0
Bld46XK	Bladder 6	16.5	0
Tst39X	Testis 1	116.6	24.35
Tst647T	Testis 2	856.16	43.5
StoAC44	Stomach 1	0	0
StoAC93	Stomach 2	0	0
SmI21XA	Small Intestine 1	68.45	0
SmIH89	Small Intestine 2	0	0
ClnAC19	Colon 1	149	21.33
ClnAS12	Colon 2	0	0
ClnB34	Colon 3	0	0
ClnB56	Colon 4	13.04	5.22
ClnAS43	Colon 5	0	0
Lng47XQ	Lung 1	0	0
Lng60XL	Lung 2	0	0
Lng75XC	Lung 3	0	3.38
Lng90X	Lung 4	0	0
LngBR26	Lung 5	0	26.82
Pan10343	Pancreas 1	50.47	0
Pan77X	Pancreas 2	281.1	0
Pan92X	Pancreas 3	18.41	0
Pan71XL	Pancreas 4	0	0
Pan82XP	Pancreas 5	0	0
PanC044	Pancreas 6	0	0
Mam12X	Mammary Gland 1	0	0
Mam162X	Mammary Gland 2	0	0
Mam42DN	Mammary Gland 3	0	0
MamS127	Mammary Gland 4	12.58	0
Mam14DN	Mammary Gland 5	0	0
End28XA	Endometrium 1	331.9	1824
	Bld46XK Tst39X Tst647T StoAC44 StoAC93 SmI21XA SmIH89 ClnAC19 ClnAS12 ClnB34 ClnB56 ClnAS43 Lng47XQ Lng60XL Lng75XC Lng90X LngBR26 Pan10343 Pan77X Pan92X Pan71XL Pan82XP PanC044 Mam12X Mam162X Mam42DN MamS127 Mam14DN	Bld46XK Bladder 6 Tst39X Testis 1 Tst647T Testis 2 StoAC44 Stomach 1 StoAC93 Stomach 2 SmI21XA Small Intestine 1 SmIH89 Small Intestine 2 ClnAC19 Colon 1 ClnAS12 Colon 2 ClnB34 Colon 3 ClnB56 Colon 4 ClnAS43 Colon 5 Lng47XQ Lung 1 Lng60XL Lung 2 Lng75XC Lung 3 Lng90X Lung 4 LngBR26 Lung 5 Pan10343 Pancreas 1 Pan77X Pancreas 2 Pan92X Pancreas 3 Pan71XL Pancreas 4 Pan82XP Pancreas 5 PanC044 Pancreas 6 Mam12X Mammary Gland 1 Mam162X Mammary Gland 3 MamS127 Mammary Gland 5	Bld46XK   Bladder 6   16.5

	End3AX	Endometrium 2	27825	65839
	End4XA	Endometrium 3	10.3	15935
	Utr1410	Uterus 1	18885	18116
	Utr23XU	Uterus 2	3358	7674
5	CvxKS52	Cervix 1	0	0
	CvxKS83	Cervix 2	0	0
	Ovr10050	Ovary 1	72.86	. <u>-</u> .
	Ovr1028	Ovary 2	0	
	Ovr638A	Ovary 3	0	
10	Ovr63A	Ovary 4	90.88	
	Ovr7730	Ovary 5	1.21	
	Ovr10400	Ovary 6	5.08	
	Ovr1050	Ovary 7	0	
	Ovr1118	Ovary 8	7.41	
15	Ovr103X	Ovary 9		32.78
	Ovr20GA	Ovary 10		0
	Ovr25GA	Ovary 11		1173.83
	Ovr35GA	Ovary 12		313.4
	Ovr50GB	Ovary 13		823.1
20	Ovr18GA	Ovary 14		40.6
	Ovr206I	Ovary 15		1264
	Ovr230A	Ovary 16		1285

0 = Negative

In the analysis of matching samples, the higher levels of expression were in prostate, endometrium, testis, and ovary showing a high degree of tissue specificity for reproductive tissues. These results confirmed the tissue specificity results obtained with the panel of normal pooled samples (Table 14).

Furthermore, the levels of mRNA expression in cancer samples and the isogenic normal adjacent tissue from the same individual were compared. This comparison provides an

indication of specificity for the cancer (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent). Table 15 shows overexpression of Prol18 in 5 out of 14 primary prostate cancer tissues (prostate samples 1, 8, 10, 11, 15) compared with their respective normal adjacent. Thus, there was overexpression in the cancer tissue for 35.71% of the prostate matching samples tested (total of 14 prostate matching samples). Expression of Prol18 was similarly higher in 3 unmatched cancer tissues (prostate samples 9, 13, 14), 10 2 prostatitis (prostate samples 20, 21), and 6 benign hyperplasia tissues (prostate samples 22 through 27).

Altogether, the high level of tissue specificity, plus the mRNA overexpression in 35.71% of the primary prostate matching samples tested are indicative of Prol18 being a diagnostic marker for prostate cancer.

- 47 -

#### What is claimed is:

1. A method for diagnosing the presence ∞ f prostate cancer in a patient comprising:

- (a) determining levels of CSG in cells, tissues or bodily 5 fluids in a patient; and
- (b) comparing the determined levels of CSG with levels of CSG in cells, tissues or bodily fluids from a normal human control, wherein a change in determined levels of CSG in said patient versus normal human control is associated with the 10 presence of prostate cancer.
  - 2. A method of diagnosing metastases of prostate cancer in a patient comprising:
  - (a) identifying a patient having prostate cancer that is not known to have metastasized;
- (b) determining CSG levels in a sample of cells, tissues, or bodily fluid from said patient; and
- (c) comparing the determined CSG levels with levels of CSG in cells, tissue, or bodily fluid of a normal human control, wherein an increase in determined CSG levels in the 20 patient versus the normal human control is associated with a cancer which has metastasized.
  - 3. A method of staging prostate cancer in a patient having prostate cancer comprising:
    - a) identifying a patient having prostate cancer;
- 25 (b) determining CSG levels in a sample of cells, tissue, or bodily fluid from said patient; and
- (c) comparing determined CSG levels with levels of CSG in cells, tissues, or bodily fluid of a normal human control, wherein an increase in determined CSG levels in said patient versus the normal human control is associated with a cancer which is progressing and a decrease in the determined CSG levels is associated with a cancer which is regressing or in remission.

4. A method of monitoring prostate cancer in a patient for the onset of metastasis comprising:

- 48 -

- (a) identifying a patient having prostate cancer that is not known to have metastasized;
- (b) periodically determining levels of CSG in samples of cells, tissues, or bodily fluid from said patient; and
- (c) comparing the periodically determined CSG levels with levels of CSG in cells, tissues, or bodily fluid of a normal human control, wherein an increase in any one of the 10 periodically determined CSG levels in the patient versus the normal human control is associated with a cancer which has metastasized.
  - 5. A method of monitoring a change in stage of prostate cancer in a patient comprising:
- (a) identifying a patient having prostate cancer;
  - (b) periodically determining levels of CSG in cells, tissues, or bodily fluid from said patient; and
- (c) comparing the periodically determined CSG levels with levels of CSG in cells, tissues, or bodily fluid of a normal human control, wherein an increase in any one of the periodically determined CSG levels in the patient versus the normal human control is associated with a cancer which is progressing in stage and a decrease is associated with a cancer which is regressing in stage or in remission.
- 6. A method of identifying potential therapeutic agents for use in imaging and treating prostate cancer comprising screening molecules for an ability to bind to CSG wherein the ability of a molecule to bind to CSG is indicative of the molecule being useful in imaging and treating prostate cancer.
- 7. The method of claim 1, 2, 3, 4, 5 or 6 wherein the CSG comprises SEQ ID NO:1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12,

- 49 -

- 13, 14, 15, 16, 17, 18, 19 or 20 or a polypeptide encoded thereby.
  - 8. An antibody which specifically binds CSG.
- A method of imaging prostate cancer in a patient
   comprising administering to the patient an antibody of claim
   8.
  - 10. The method of claim 9 wherein said antibody is labeled with paramagnetic ions or a radioisotope.
- 11. A method of treating prostate cancer in a patient 10 comprising administering to the patient an antibody of claim 7.
  - 12. The method of claim 11 wherein the antibody is conjugated to a cytotoxic agent.

#### SEQUENCE LISTING

```
<110> Salceda, Susana
        Recipon, Herve
        Cafferkey, Robert
        diaDexus, LLC
  <120> Method of Diagnosing, Monitoring, Staging, Imaging and
        Treating Prostate Cancer
  <130> DEX-0052
  <140>
  <141>
 <150> 60/104,737
  <151> 1998-10-19
 <160> 36
 <170> PatentIn Ver. 2.0
 <210> 1
 <211> 188
 <212> DNA
 <213> Homo sapiens
 <400> 1
ggtaaacacc tgcttttatc atcagaacaa agaggctgtg tcccctgccc tatgaggtcc 60
atttctgaga gttgtggcta atgggcaaga aggttggggc tttagagatt tgggataaag 120
atatcaaaca ccagaaaggt agaaagaagt gatcagatta gggttactta ggtgatgata 180
tgaactct
<210> 2
<211> 9819
<212> DNA
<213> Homo sapiens
<400> 2
cagetggggt ctacccaggt ccatgtettg gacatgttga gagtttttet ggaaggcagg 60
gatacagtgt ggtccaaaaa cacacaaatg cccctactgg cccaggggtt gtcacaatag 120
actggaaggg tgacacatcc caggcgcttg ccacccatca cacgcacctc ctacccactg 180
gcatccttcc accccaggca cacacaaagc ctcagtccag agatcaactc tggactcagc 240
tetgaatttg catateetgt gtgtagatte attetteata acetetgeee ageetagett 300
gtgtatcatt tttttttctc tattagggga ggagcccgtc ctggcactcc cattggcctg 360
tagattcacc tcccctgggc agggccccag gacccaggat aatatctgtg cctcctgccc 420
agaaccetee aageagacae aatggtaaga atggtgeetg teetgetgte tetgetgetg 480
cttctgggtc ctgctgtccc ccaggagaac caagatggtg agtggggaaa gcaagggatg 540
```

ggtgctgga	ng aggactgga	a ggaggtgag	g aacaggaca	t gtggctggg	a gacaggetge	g 600
atgcagctg	g gataccctg	g catacggca	g gaatgggtg	c ccaaggctg	t caactccct	660
	a cttccagga					
	ga atctaaaac					
	g ggtagacag				All and a second	
	t teceteette					
	c aaggggaaag					
	c aaatcccaca					
	a gagtgggag					
	g gcatgagaat					
	a aaattcatga					
	a ctttggaatg					
	a acagaactaa					
	c tgtatacaca					
	g agagagagta					
	c ctaaatcttc					
	t tgattccaag					
	t ttctcttaag					
	t actcaaggtc					
	g tatatattta					
	a tcatcaaaaa					
	a attatactt					
	a ttttttgagac					
	cactgcaacc					
	aggactacag					
	g tttcatcatg					
	ctcccaaagt					
	taaaagtgta					
	tcacccaggc					
	ttcaagcaat					
	aatagagata					
	tccactcact					
	ccagttaaat					
	cattettett					
	aatcttggct					
	ctgagtcgct					
	agagatgggg					
	acctgcctca					
	gttaaattat					
	tttttgtacc					
	cctttggtaa					
	ttaaaggcta					
	attctgccat					
	atggaactat					
	tacaaccccg					
	catttcttga					
	aggttaagct					
tttgtcacct	aggctgaagt	gcagtggcat	gatctcagct	cactgcaagc	tccgcctccc	3420

gggttcatgc cattetectg ceteageete etgagtaget gggaetaeag geaceegeea 3480 ccatgcttgg ctaatttttt gaattcttag tagagacggg gtttcaccat gttagccagg 3540 atggtctcga tctcctgacc tcgtgatcca cccgcctcgg ccccctaaag tgctgggatt 3600 acaggcgtga gccactgagc ccggacgaaa tgttaatttg tttttttga gacggagtct 3660 cactetgtea tecaagetgg agtgeagtgg catgatettg gettgttgea acctetgeet 3720 ctctggttca agtgattttc ctgcctcagc ctccagcatg actgggatta caggcccgca 3780 ccaccatgcc cagctaattt ttgtattttt taatagagat ggggtttcac catgttggcc 3840 aggetggtet teaacteetg ateteaagta atetgeetge ettggeetee caaagteetg 3900 ggattacagg catgagccac ggagcccagc ctagaaatgt taatttctaa cgcatgtcag 3960 attccatgca cactgggcaa ggttccattc ctccatgggg tgactcaggg atccaggcca 4020 attgcatatt gagactettt catattatee tgtggeette aaagtegtea eetetaggga 4080 tgagaaacaa aagggaaagc cagctggtag ggtcttggac aagaagaaag acatcacttc 4140 tgctcacatt ctcttttgac aaaactcagt cacatggtcc caatatatct tcgaggtggc 4200 tgagtaatgt tatetteeta tgtgteaage agaggaaata atgtagtgaa gacacaggat 4260 ggtctctgaa atatcatctc aggcatgaaa gtagagcata ttcacttgag tgagcctcca 4320 gtggtgtgaa gttgatggca ggagaaagag ctggggaaga aaaggccagt ggcaggtctc 4380 ccctcctagc cctatgcagc cccacagtgg gacccttgca tggacctcaa ccatcagaat 4440 cttttctttt gcaggtcgtt actctctgac ctatatctac actgggctgt ccaagcatgt 4500 tgaagacgtc cccgcgtttc aggcccttgg ctcactcaat gacctccagt tctttagata 4560 caacagtaaa gacaggaagt ctcagcccat gggactctgg agacaggtgg aaggaatgga 4620 ggattggaag caggacagcc aacttcagaa ggccagggag gacatcttta tggagaccct 4680 gaaagacatt gtggagtatt acaacgacag taacggtcag tgaataacag accacagggg 4740 tggaaggtct aacccaagag gcagccccc cagtgtgagt ggcaagggat cagcaggatg 4800 gaaatagtee caateecagg ggaagaacag gagacacage agaaacacag acatgteege 4860 ateccaecca ecceacagea caggtgetee eegetteece ateaattgee ecatecteat 4920 cccaggeete aggteaeaea ggaagtgatg geagagteae tteetateea ggeaeetatg 4980 accteteace tecacacece acceategga ggetgatace ecegtgagaa ggeateagae 5040 tracccctgt ccagggaggt tgcctggaga gtgagccact ctcaaagtca ctcagacctg 5100 ggctcacctg gtggttctgc cagtcctagc tgttgacagt gaaacgttcc caaaatatct 5160 ggttgaaatc tgcaaacatt ggagcactga gacctacctc caaacaagtc tgtaatattt 5220 aactatgtet gttetatgaa ggatgteaca gtetgteetg ateteeettg eageteeate 5280 acctagcaca gggtacagcc aatattggct caattgaaat ttgtggaatc cacagagaaa 5340 agcacccggc acacaccgta gcccatgctg ggggctcagg aagtgctgga ttcaaaactg 5400 tgggctgtta gagttccttg gagccctaaa gttcctcctt accatacgat gcagacccag 5460 gaagggccac ctgcgctatg gtcagaggag ctggtggcag agcccgtgca gagatggtcc 5520 etgtgeeece ggeecagtge tettteteet aaaccacact geeageeca aggeageeaa 5580 cctcaggtct ggtgaactgc tggtgttaaa ttatcataga gtgggtgtca aaagatgggc 5640 tactaagtac aaaaatgccc aaggtgctac atgggatctg aagattttca aaaggaggca 5700 ggcccaggct gtgtgtcagc aataggagag gagggggcac aggtgatcag aaaagacact 5820 gggggaagca ttgatggaca ggaatagaaa tggcaaagtg gataattaag aggaaggagg 5880 atgaggagat gaacacaggg tattagaaaa taatagaagg cagggettgg tggetcacte 5940 ttgtaateee ageaetttgg gaggetgagg caggeagate acctaaggte aggagttega 6000 gaccageceg gecaacatgg tgaaaccetg tetetactaa taatacaaaa atageetgge 6060 atggtggcac acgtctgtgg tcccagctac tcaggaggct gaggcaggag aattgcttga 6120 acccaggagg cagaggttac agtggccaaa atcctaccat tgcactacag cctgggtgac 6180 aagagtgaaa cgttgtctaa aaacaaaaaa caaaaaacaa aaaaaggaaa taatagtagc 6240 tgacatttac tgagcactta ctttgtgcca ggcccatcta tgagcatata taatgctcag 6300

aatagcccc	c taaaacagt	g ctcttggcat	tgccatttca	a gaggtgagga	a aatagaggca	6360
cagggagtt	g agtggctcca	a gttcaggcaa	a cacaccaggt	gggggtggg	g ggctggggag	6420
agacctggg	a cgtgagccca	a gacagettga	a gagettteag	g agtctatgc	aacagcacca	6480
accagtgct	g ggtaaacac	tgcttttatc	atcagaacaa	agaggctgtg	g teceetgeed	6540
tatgaggtc	c atttctgaga	a gttgtggcta	a atgggcaaga	aggttgggg	tttagagatt	6600
tgggataaag	g atatcaaaca	a ccagaaaggt	agaaagaagt	gatcagatta	a gggttactta	6660
ggtgatgata	a tgaactctto	ctagaactga	a gagaaaaaga	gageetteet	ttactcatat	6720
gaaatcacaa	a ataatttcta	tccaatttgg	g aagtacactt	tggtgtagtt	gtgacagett	6780
cctcaggact	cagcataaat	tcaaacaaat	aattgtcctt	agaagagat	, ctatagaaga	6840
gatagaaata	a tattcatatt	ctgtagcttt	ttttttttg	agatggagtt	: ttgctcttgt	6900
cacccaagct	ggagtgcagt	gatgcaatct	cagctcactg	caaactttgc	ctcctgggtt:	6960
caagggatto	tcctgcctca	gcctcccgat	aactgggact	acaggetaca	ggcatgtgtc	7020
actactcct	gttaatttt	tttttttt	tttaagactg	agtettgete	tgtctttcag	7080
gctgatgtac	: aatggctcca	tctcggctca	ctacaacttc	tgtcccccag	gttcaagcga	7140
tteteetgee	: tcagcctcat	gagtagctgg	gattacaggc	atgtgccago	acacccagca	7200
aatttttgta	tttttagtag	agatgaggtc	ttaccatgtt	ggccaggctg	gtctcaaact	7260
cctgacctca	ggtgatcctt	tggcctcagc	ctccctaact	gctgggatta	caggcatgag	7320
ccactgcgtc	cagectaatt	ttatatttt	ggtagagatg	gggtttcacc	atattggcca	7380
ggctggtctc	gaactcatga	cctaaggtga	tccatcctcc	tcagcctctc	aaagtgctgg	7440
gattacaagt	gtgagccact	gggcctggtg	ctttttttt	tttttttt	tttttttt	7500
tgagataggg	tctcactctg	tcacccaggc	tgaaatgcag	tagtgtgatt	ttggctcatt	7560
gcagccttga	cttcccaggc	tgaagtgatc	ctcccacctc	agcctcctga	gtagctgggg	7620
ctacaggcat	gcaccaccat	gctgcgctaa	tttttatatt	ttttgtagtg	gtgggatttc	7680
gccatatcac	cctggctggt	ctggaacccc	tgggctcaag	cgatccactc	gcttcagctt	7740
ctcaaagtgc	tgggattaca	ggcatgagcc	acagcgccca	ggctgtagct	ctcttaagga	7800
ggaacatatc	tcatctgaga	caaacctgaa	atgccaaacc	aaactgagtt	agcccctctc	7860
tgtctgttgt	atatattgga	gtaataacct	atttgtcttg	ataaagggat	tgcatgcttg	7920
aattgcaaaa	acctttattt	cttttgggtt	gcccaatgtg	caagactaag	agttattttg	7980
ataaatttct	caccaggctg	actgtctctc	tgtggggtcg	ggggagtttt	cagggtctca	8040
cgtattgcag	ggaaggtttg	gttgtgagat	cgagaataac	agaagcagcg	gagcattctg	8100
gaaatattac	tatgatggaa	aggactacat	tgaattcaac	aaagaaatcc	cagcctgggt	8160
ccccttcgac	ccagcagccc	agataaccaa	gcagaagtgg	gaggcagaac	cagtctacgt	8220
gcagcgggcc	aaggcttacc	tggaggagga	gtgccctgcg	actctgcgga	aatacctgaa	8280
atacagcaaa	aatatcctgg	accggcaagg	tactcactgc	ttcctgctcc	ccagtactga	8340
gcccagaata	aaagacgatc	tcaggctagg	agctcaggca	acatcttagt	ccggtctcat	8400
ctgttcctgg	atgtccctca	gacccccagc	tttcatcttt	taggatttat	tccttccctg	8460
ggataatata	atttgtggtc	caaaaagaac	atcatcaaaa	tttcaggcag	aatgggccag	8520
gaaggccatt	ctttcttgat	gagtgtcccc	aaatcatctc	caattaacag	acaaggagct	8580
tgaggttagg	gaggtgaggg	taacactgtc	tgtaagaggc	agagctggga	ctcaaattcc	8640
agatttcaga	ttccaaatcc	catcgttttt	tatctctaca	atgatgcctc	ccatctgggt	8700
					agccagtgtg	
agcggaattg	atggctgcaa	gctgagactt	ggattggaga	cgtagtgaga	ctcaggattg	8820
					gcagctggac	
					gagaagcaca	
					ggattgggca	
					gggggtgggc	
					aggaagagac	
ctgggccggg	agaagtccac	ctcaagcctg	cagtgtcaca	ctctatccct	ccacagatcc	9180

```
tecetetgtg gtggteacea gecaceagge eccaggagaa aagaagaaac tgaagtgeet 9240
  ggcctacgac ttctacccag ggaaaattga tgtgcactgg actcgggccg gcgaggtgca 9300
  ggagcctgag ttacggggag atgttcttca caatggaaat ggcacttacc agtcctgggt 9360
  ggtggtggca gtgccccgc aggacacagc cccctactcc tgccacgtgc agcacagcag 9420
  cctggcccag cccctcgtgg tgccctggga ggccagctag gaagcaaggg ttggaggcaa 9480
  tgtgggatct cagacccagt agctgccctt cctgcctgat gtgggagctg aaccacagaa 9540
  atcacagtca atggatccac aaggeetgag gageagtgtg gggggacaga caggaggtgg 9600
  atttggagac cgaagactgg gatgcctgtc ttgagtagac ttggacccaa aaaatcatct 9660
  caccttgage ccacceccae eccattgtet aatetgtaga agetaataaa taateateee 9720
  teettgeeta geataacaga gaateetttt tttaaeggtg atgegetgta gaaatgtgae 9780
  tagattttct cattggttct gccctcaagc actgaattc
                                                                     9819
  <210> 3
  <211> 250
  <212> DNA
  <213> Homo sapiens
 <400> 3
 cgcccctgcg ccgccgagcc agctgccaga atgccgaact ggggaggagg caagaaatgt 60
 ggggtgtgtc agaagacggt ttactttgcc gaagaggttc agtgcgaagg caacagcttc 120
 cataaatcct gcttcctgtg catggtctgc aagaagaatc tggacagtac cactgtggcc 180
 gtgcatggtg aggagattta ctgcaagtcc tgctacggca agaagtatgg gcccaaaggc 240
 tatggctacg
                                                                    250
 <210> 4
 <211> 1900
 <212> DNA
 <213> Homo sapiens
 <220>
 <221> unsure
 <222> (16)
<220>
<221> unsure
<222> (18)
<220>
<221> unsure
<222> (20)
<220>
<221> unsure
<222> (1887)
<220>
<221> unsure
<222> (1894)
```

```
<400> 4
 acgccttccg cggagnanan caaaacggcg cgcaggccgg gcgcacccag ccgccacttc 60
 cgagagegee tgeegeeet ggegeegeeg agecagetge cagaatgeeg aactggggag 120
 gaggcaagaa atgtggggtg tgtcaagaag acggtttact ttgccgaaga ggttcagtgc 180
 gaaggcaaca gcttccataa atcctgcttc ctgtgcatgg tctgcaagaa gaatctggac 240
 agtaccactg tgggccgtgc atggtgagga gatttactgg caagtccctg ctacggcaag 300
 aagtatgggc ccaaaggcta tggctacggg ccagggcgca ggcaccctca gcactgacaa 360
 gggggagtcg ctgggtatca agcacgagga agcccctggg ccacaggccc accaccaacc 420
 ccaatggcat ccaaatttgc ccagaagatt ggtggctccg agegctgccc ccgatgcagc 480
 caggcagtct atgctgcgga gaaggtgatt ggtgctggga agtcctggca taaggcctgc 540
 tttcgatgtg ccaagtgtgg caaaggcctt gagtcaacca ccctgggcag acaaggatgg 600
 cgagatttac tgcaaaggat gttatgctaa aaacttcggg cccaagggct ttggttttgg 660
 gcaaggagct ggggccttgg tccactctga gtgaggccac catcacccac cacaccctgc 720
 ccactcctgc gcttttcatc gccattccat tcccagcagc tttggagacc tccaggatta 780
 tttctctgtc agccctgcca catatcacta atgacttgaa cttgggcatc tggctccctt 840
 tggtttgggg gtctgcctga ggtcccaccc cactaaaggg ctccccaggc ctgggatctg 900
 acaccatcac cagtaggaga cctcagtgtt ttgggtctag gtgagagcag gcccctctcc 960
ccacacctcg ccccacagag ctctgttctt agcctcctgt gctgcgtgtc catcatcagc 1020
tgaccaagac acctgaggac acatettggc acceagagga gcagcagcaa caggetggag 1080
ggagagggaa gcaagaccaa gatgaggagg ggggaaggct gggttttttg gatctcagag 1140
atteteetet gtgggaaaga ggttgagett cetggtgtee eteagagtaa geetgaggag 1200
teccagetta gggagtteae tattggagge agagaggeat geaggeaggg tectaggage 1260
ccctgcttct ccaggcctct tgcctttgag tctttgtgga atggatagcc tcccactagg 1320
actgggagga gaataaccca ggtcttaagg accccaaagt caggatgttg tttgatcttc 1380
tcaaacatct agttccctgc ttgatgggag gatcctaatg aaatacctga aacatatatt 1440
ggcatttatc aatggctcaa atcttcattt atctctggcc ttaaccctgg ctcctgaggc 1500
tgcggccagc agagcccagg ccagggctct gttcttgcca cacctgcttg atcctcagat 1560
gtggagggag gtaggcactg cctcagtctt catccaaaca cctttccctt tgccctgaga 1620
cctcagaatc ttccctttaa cccaagaccc tgcctcttcc actccaccct tctccaggga 1680
cccttagatc acatcactcc acccctgcca ggccccaggt taggaatagt ggtgggagga 1740
aggggaaagg gctgggcctc accgctccca gcaactgaaa ggacaacact atctggagcc 1800
acceaetgaa agggetgeag geatgggetg tacceaaget gattteteat etggteaata 1860
aagctgttta gaccagaaaa aaaaaanaaa aaanaaaagg
                                                                  1900
<210> 5
<211> 273
<212> DNA
<213> Homo sapiens
<400> 5
gatgcatcaa aagagctgca agttctccac attgacttct tgaatcagga caacgccgtt 60
tctcaccaca catgggagtt ccaaacgagc agtcctgtgt tccggcgagg acaggtgttt 120
cacctgcggc tggtgctgaa ccagccccta caatcctacc accaactgaa actggaattc 180
agcacagggc cgaatcctag catcgccaaa cacaccctgg tggtgctcga cccgaggacg 240
ccctcagacc actacaactg gcaggcaacc ctt
                                                                  273
<210> 6
```

6

<211> 3021 <212> DNA <213> Homo sapiens

<400> 6 tgtggaagca ccaggcatca gagatagagt cttccctggc attgcaggag agaatctgaa 60 gggatgatgg atgcatcaaa agagctgcaa gttctccaca ttgacttctt gaatcaggac 120 aacgccgttt ctcaccacac atgggagttc caaacgagca gtcctgtgtt ccggcgagga 180 caggtgtttc acctgcggct ggtgctgaac cagcccctac aatcctacca ccaactgaaa 240 ctggaattca gcacagggcc gaatcctagc atcgccaaac acaccctggt ggtgctcgac 300 ccgaggacgc cctcagacca ctacaactgg caggcaaccc ttcaaaatga gtctggcaaa 360 gaggtcacag tggctgtcac cagttccccc aatgccatcc tgggcaagta ccaactaaac 420 gtgaaaactg gaaaccacat cettaagtet gaagaaaaca teetataeet tetetteaac 480 ccatggtgta aagaggacat ggttttcatg cctgatgagg acgagcgcaa agagtacatc 540 ctcaatgaca cgggctgcca ttacgtgggg gctgccagaa gtatcaaatg caaaccctgg 600 aactttggtc agtttgagaa aaatgtcctg gactgctgca tttccctgct gactgagagc 660 teceteaage ceacagatag gagggaeeee gtgetggtgt geagggeeat gtgtgetatg 720 atgagetttg agaaaggeea gggegtgete attgggaatt ggaetgggga etatgaaggt 780 ggcacagece catacaagtg gacaggcagt geeegatee tgcageagta etacaacaeg 840 aagcaggetg tgtgetttgg ccagtgetgg gtgtttgetg ggateetgae tacagtgetg 900 agagegttgg geateceage aegeagtgtg acaggetteg atteagetea egacacagaa 960 aggaacctca cggtggacac ctatgtgaat gagaatggca agaaaatcac cagtatgacc 1020 cacgactetg tetggaattt ceatgtgtgg acggatgeet ggatgaageg aceggatetg 1080 cccaagggct acgacggctg gcaggctgtg gacgcaacgc cgcaggagcg aagccagggt 1140 gtettetget gtgggecate accaetgace gecateegea aaggtgacat etttattgte 1200 tatgacacca gattcgtctt ctcagaagtg aatggtgaca ggctcatctg gttggtgaag 1260 atggtgaatg ggcaggagga gttacacgta atttcaatgg agaccacaag catcgggaaa 1320 aacatcagca ccaaggcagt gggccaagac aggcggagag atatcaccta tgagtacaag 1380 tatccagaag geteetetga ggagaggeag gtteatggat catgeettee teetteteag 1440 ttctgagagg gagcacagac gacctgtaaa agagaacttt cttcacatgt cggtacaatc 1500 agatgatgtg ctgctgggaa actctgttaa tttcaccgtg attcttaaaa ggaagaccgc 1560 tgccctacag aatgtcaaca tcttgggctc ctttgaacta cagttgtaca ctggcaagaa 1620 gatggcaaaa ctgtgtgacc tcaataagac ctcgcagatc caaggtcaag tatcagaagt 1680 gactetgace ttggaeteca agacetacat caacageetg getatattag atgatgagee 1740 agttatcaga ggtttcatca ttgcggaaat tgtggagtct aaggaaatca tggcctctga 1800 agtattcacg tetttecagt accetgagtt etetatagag ttgeetaaca caggeagaat 1860 tggccagcta cttgtctgca attgtatctt caagaatacc ctggccatcc ccttgactga 1920 cgtcaagttc tctttggaaa gcctgggcat ctcctcacta cagacctctg accatgggtg 1980 agtotgootg aggaoggtgo agootggtga gaccatocaa toocaaataa aatgoaccoo 2040 aatgeteaga agattgttet cateaceaag tageettgte tgatgetgtg gageettagt 2160 tgagatttca gcatttccta ccttgtggct tagctttcag attatggatg attaaatttg 2220 atgacttata tgagggcaga ttcaagagcc agcaggtcaa aaaggccaac acaaccataa 2280 gcagccagac ccacaaggcc aggtcctgtg ctatcacagg gtcaccttct tttacagtta 2340 gaaacaccag ccgaggccac agaatcccat ccctttcctg agtcatggcc tcaaaaatca 2400 gggccaccat tgtctcaatt caaatccata gatttcgaag ccacagattc tctccctgga 2460 gcaagcatga ctatgggcag cccagtgctg ccacctgctg acgacccttg agaagctgcc 2520 atatetteag gecatgggtt caccageeet gaaggeaeet gteaaetgga gtgetetete 2580

```
agcactggga tgggcctgat agaagtgcat tetectecta ttgcctccat tetectetet 2640
 ctatccctga aatccaggaa gtccctctcc tggtgctcca agcagtttga agcccaatct 2700
 gcaaggacat ttctcaaggg ccatgtggtt ttgcagacaa ccctgtcctc aggcctgaac 2760
 tcaccataga gacccatgtc agcaaacggt gaccagcaaa tcctcttccc ttattctaaa 2820
 gctgcccctt gggagactcc agggagaagg cattgcttcc tccctggtgt gaactctttc 2880
 tttggtattc catccactat cctggcaact caaggctgct tctgttaact gaagcctgct 2940
 cettettgtt etgeceteca gagatttget caaatgatea ataagettta aattaaacte 3000
 tacttcaaga aaaaaaaacc g
 <210> 7
 <211> 267
 <212> DNA
 <213> Homo sapiens
 <400> 7
gaacattcca gatacctatc attactcgat gctgttgata acagcaagat ggctttgaac 60 -
tcagggtcac caccagctat tggaccttac tatgaaaacc atggatacca accggaaaac 120
ccctatcccg cacagcccac tgtggtcccc actgtctacg aggtgcatcc ggctcagtac 180
tacccgtccc ccgtgcccca gtacgccccg agggtcctga cgcaggcttc caaccccgtc 240
gtctgcacgc agcccaaatc cccatcc
                                                                   267
<210> 8
<211> 3443
<212> DNA
<213> Homo sapiens
<400> 8
gggcgggccg ggccgagtag gcgcgagcta agcaggaggc ggaggcggag gcggagggcg 60
aggggcgggg agcgccgcct ggagcgcggc aggtcatatt gaacattcca gatacctatc 120
attactogat gotgttgata acagcaagat ggotttgaac toagggtcac caccagotat 180
tggaccttac tatgaaaacc atggatacca accggaaaac ccctatcccg cacagcccac 240
tgtggtcccc actgtctacg aggtgcatcc ggctcagtac tacccgtccc ccgtgcccca 300
gtacgccccg agggtcctga cgcaggcttc caaccccgtc gtctgcacgc agcccaaatc 360
cccatccggg acagtgtgca cctcaaagac taagaaagca ctgtgcatca ccttgaccct 420
ggggaccttc ctcgtgggag ctgcgctggc cgctggccta ctctggaagt tcatgggcag 480
caagtgctcc aactctggga tagagtgcga ctcctcaggt acctgcatca acccctctaa 540
ctggtgtgat ggcgtgtcac actgccccgg cggggaggac gagaatcggt gtgttcgcct 600
ctacggacca aacttcatcc ttcaggtgta ctcatctcag aggaagtcct ggcaccctgt 660
gtgccaagac gactggaacg agaactacgg gcgggcggcc tgcagggaca tgggctataa 720
gaataatttt tactctagcc aaggaatagt ggatgacagc ggatccacca gctttatgaa 780
actgaacaca agtgccggca atgtcgatat ctataaaaaa ctgtaccaca gtgatgcctg 840
ttcttcaaaa gcagtggttt ctttacgctg tatagcctgc ggggtcaact tgaactcaag 900
ccgccagagc aggatcgtgg gcggcgagag cgcgctcccg ggggcctggc cctgggcagg 960
tcagcctgca cgtccagaac gtccacgtgt gcggaggctc catcatcacc cccgagtgga 1020
tcgtgacagc cgcccactgc gtggaaaaac ctcttaacaa tccatggcat tggacggcat 1080
ttgcggggat tttgagacaa tctttcatgt tctatggagc cggataccaa gtagaaaaag 1140
tgatttctca tccaaattat gactccaaga ccaagaacaa tgacattgcg ctgatgaagc 1200
tgcagaagcc tctgactttc aacgacctag tgaaaccagt gtgtctgccc aacccaggca 1260
```

```
tgatgctgca gccagaacag ctctgctgga tttccgggtg gggggccacc gaggagaaag 1320
  ggaagacete agaagtgetg aacgetgeea aggtgettet cattgagaca cagagatgea 1380
  acagcagata tgtctatgac aacctgatca caccagccat gatctgtgcc ggcttcctgc 1440
  aggggaacgt cgattettge cagggtgaca gtggagggcc tetggteact teggaggaaca 1500
  atatctggtg gctgataggg gatacaagct ggggttctgg ctgtgccaaa gcttacagac 1560
  caggagtgta cgggaatgtg atggtattca cggactggat ttatcgacaa atgagggcag 1620
  acggctaatc cacatggtct tegteettga egtegtttta caagaaaaca atggggetgg 1680
  ttttgcttcc ccgtgcatga tttactctta gagatgattc agaggtcact tcatttttat 1740
  taaacagtga acttgtctgg ctttggcact ctctgccatt ctgtgcaggc tgcagtggct 1800
  cccctgccca gcctgctctc cctaacccct tgtccgcaag gggtgatggc cggctggttg 1860
  tgggcactgg cggtcaagtg tggaggagag gggtggaggc tgccccattg agatcttcct 1920
  gctgagtcct ttccaggggc caattttgga tgagcatgga gctgtcacct ctcagctgct 1980
  ggatgacttg agatgaaaaa ggagagacat ggaaagggag acagccaggt ggcacctgca 2040
  geggetgeet etggggeeae ttggtagtgt ecceageeta eeteteeaea aggggatttt 2100
  gctgatgggt tcttagagcc ttagcagccc tggatggtgg ccagaaataa agggaccagc 2160
  ccttcatggg tggtgacgtg gtagtcacct tgtaagggga acagaaacat ttttgttctt 2220
  atggggtgag aatatagaca gtgcccttgg gtgcgaggga agcaattgaa aaggaacttg 2280
 ccctgagcac tcctggtgca ggtctccacc tgcacattgg gtggggctcc tgggagggag 2340
 actcagcctt cctcctcatc ctccctgacc ctgctcctag caccctggag agtgcacatg 2400
 ccccttggtc ctgggcaggg gcgccaagtc tggcaccatg ttggcctctt caggcctgct 2460
 agtcactgga aattgaggtc catgggggaa atcaaggatg ctcagtttaa ggtacactgt 2520
 ttccatgtta tgtttctaca cattgctacc tcagtgctcc tggaaactta gcttttgatg 2580
 tetecaagta gtecaeette atttaaetet ttgaaaetgt ateatetttg eeaagtaaga 2640
 gtggtggcct atttcagctg ctttgacaaa atgactggct cctgacttaa cgttctataa 2700
 atgaatgtgc tgaagcaaag tgcccatggt ggcggcgaag aagagaaaga tgtgttttgt 2760
 tttggactet etgtggteee ttecaatget gtgggtttee aaccagggga agggteeett 2820
 ttgcattgcc aagtgccata accatgagca ctactctacc atggttctgc ctcctggcca 2880
 agcaggctgg tttgcaagaa tgaaatgaat gattctacag ctaggactta accttgaaat 2940
 ggaaagtett geaateeeat ttgeaggate egtetgtgea eatgeetetg tagagageag 3000
 catteccagg gacettggaa acagttggea etgtaaggtg ettgeteece aagacacate 3060
 ctaaaaggtg ttgtaatggt gaaaacgtct tccttcttta ttgccccttc ttatttatgt 3120
gaacaactgt ttgtcttttt ttgtatcttt tttaaactgt aaagttcaat tgtgaaaatg 3180
aatatcatgc aaataaatta tgcgattttt ttttcaaagt aaccactgca tctttgaagt 3240
tetgeetggt gagtaggace ageeteeatt teettataag ggggtgatgt tgaggetget 3300
ggtcagagga ccaaaggtga ggcaaggcca gacttggtgc tcctgtggtt ggtgccctca 3360
gttcctgcag cctgtcctgt tggagaggtc cctcaaatga ctccttctta ttattctatt 3420
agtctgtttc catgggcgtg ata
                                                                   3443
<210> 9
<211> 254
<212> DNA
<213> Homo sapiens
<400> 9
gtgctgcacc aggccaccat cctgcccaag actgggacag tgtccctgga ggtacggctc 60
ctggaggcct cccgtgcctt cgaggtgtca gagaacggca acctggtagt gagtgggaag 120
gtgtaccagt gggatgaccc tgaccccagg ctcttcgacc acccggaaag ccccaccccc 180
aaccccacgg agcccctctt cctggcccag gctgaagttt acaaggagct gcgtctgcgt 240
```

```
ggctacgact acgg
                                                                    254
 <210> 10
 <211> 8470
 <212> DNA
 <213> Homo sapiens
 <220>
 <221> unsure
 <222> (4131)
 <220>
 <221> unsure
 <222> (5117)
 <220>
 <221> unsure
 <222> (5552)
<400> 10
cggccgtcga cacggcagcg gccccggcct ccctctccgc cgcgcttcag cctcccgctc 60
cgccgcgctc cagcctcgct ctccgccgcc cgcaccgccg cccgcgccct caccagagca 120
gccatggagg aggtggtgat tgccggcatg tccgggaagc tgccagagtc ggagaacttg 180
caggagttct gggacaacct catcggcggt gtggacatgg tcacggacga tgaccgtcgc 240
tggaaggcgg ggctctacgg cctgccccgg cggtccggca agctgaagga cctgtctagg 300
tttgatgcct ccttcttcgg agtccacccc aagcaggcac acacgatgga ccctcagctg 360
eggetgetge tggaagteae etatgaagee ategtggaeg gaggeateaa eccagattea 420
ctccgaggaa cacacactgg cgtctgggtg ggcgtgagcg gctctgagac ctcggaggcc 480
ctgagccgag accccgagac actcgtgggc tacagcatgg tgggctgcca gcgagcgatg 540
atggccaacc ggctctcctt cttcttcgac ttcagagggc ccagcatcgc actggacaca 600
geetgeteet ceageetgat ggeeetgeag aaegeetaee aggeeateea cagegggeag 660
tgccctgccg ccatcgtggg gggcatcaat gtcctgctga agcccaacac ctccgtgcag 720
ttcttgaggc tggggatgct cagccccgag ggcacctgca aggccttcga cacagcgggg 780
aatgggtact gccgctcgga gggtgtggtg gccgtcctgc tgaccaagaa gtccctggcc 840
eggegggtgt aegecaceat eetgaaegee ggeaecaata eagatggett eaaggageaa 900
ggcgtgacct tcccctcagg ggatatccag gagcagctca tccgctcgtt gtaccagtcg 960
gccggagtgg cccctgagtc atttgaatac atcgaagccc acggcacagg caccaaggtg 1020
ggcgaccccc aggagctgaa tggcatcacc cgagccctgt gcgccacccg ccaggagccg 1080
ctgctcatcg gctccaccaa gtccaacatg gggcaccegg agccagecte ggggctggca 1140
gccctggcca aggtgctgct gtccctggag cacgggctct gggcccccaa cctgcacttc 1200
catagececa accetgagat eccagegetg ttggatggge ggetgeaggt ggtggaceag 1260
cccctgcccg tccgtggcgg caacgtgggc atcaactcct ttggcttcgg gggctccaaa 1320
egtgeacate atectgagge ceaacaegea geogeeeeee geaceeggee cacatgeeae 1380
cctgccccgt ctgctgcggg ccagcggacg cacccctgag gccgtgcaga agctgctgga 1440
gcagggcctc cggcacagcc agggcctggc tttcctgagc atgtgaacga catcgcggct 1500
gtccccgacc accgccatgc ccttccgtgg ctacgctgtg ctgggtggtg agacgcggtg 1560
geceagaggt geageaggtg eeegetggeg agegeeeget etggtteate tgetetggga 1620
tgggcacaca gtggcgcggg atggggctga gcctcatgcg cctggaccgc ttccgagatt 1680
```

CCATCCT and otherwise gotter
ccatcctacg ctccgatgag gctgtgaacc gattcggcct gaaggtgtca cagctgctgc 1740
tgagcacaga cgagagcacc tttgatgaca tcgtccattc gtttgtgagc ctgactgcca 1800
tecagatagg ceteatagae etgetgaget geatgggget gaggeeagat ggeategteg 1860
gccactccct gggggaggtg gcctgtggct acgccgacgg ctgcctgtcc caggaggagg 1920
ccgtcctcgc tgcctactgg aggggacagt gcatcaaaga agcccatctc ccgccgggcg 1980
ccatggcagc cgtgggcttg tcctgggagg agtgtaaaca gcgctgcccc ccggcggtgg 2040
tgcccgccgc cacaactcca aggacacagt caccatctcg ggacctcagg ccccggtgtt 2100
tgagttcgtg gagcagctga ggaaggaggg tgtgtttgcc aaggaggtgc ggaccggcgg 2160
tatggcette cactectaet teatggagge categeacee ceaetgetge aggageteaa 2220
gaaggtgate egggageega agecaegtte agecegetgg eteageaeet etateeeega 2280
ggcccagtgg cacagcagcc tggcacgcac gtcctccgcc gagtacaatg tcaacaacct 2340
ggtgageeet gtgetgttee aggaggeeet gtggeaegtg eetgageaeg eggtggtget 2400
ggagatcgcg ccccacgccc tgctgcaggc tgtcctgaag cgtggcctga agccgagctg 2460
caccatcate eccetgatga agaaggatea cagggacaae etggagttet teetggeegg 2520
categgeagg etgeacetet eaggeatega egecaacece aatgeettgt teccacetgt 2580
ggagtcccca gctccccgag gaactcccct catctcccca ctcatcaagt gggaccacag 2640
cctggcctgg gacgcgcgg ccgccgagga cttccccaac ggttcaggtt ccccctcagc 2700
caccatctac acatgcacac caagctccga gtctcctgac cgctacctgg tggaccacac 2760
categaeggt egegteetet teecegeeae tggetaeetg ageatagtgt ggaagaeget 2820
ggcccgaccc ctgggcctgg gcgtcgagca gctgcctgtg gtgtttgagg atgtggtgct 2880
gcaccaggcc accatectgc ccaagactgg gacagtgtcc ctggaggtac ggctcctgga 2940
ggcctcccgt gccttcgagg tgtcagagaa cggcaacctg gtagtgagtg ggaaggtgta 3000
ccagtgggat gaccctgacc ccaggctctt cgaccacccg gaaagcccca cccccaaccc 3060
cacggageee etetteetgg eccaggetga agtttacaag gagetgegte tgegtggeta 3120
cgactacggc cctcatttcc agggcatcct ggaggccagc ctggaaggtg actcggggag 3180
gctgctgtgg aaggataatg ggtgagttca tggacaccat gctgcagatg tccatcctgg 3240
gteggecaag caeggeetgt acetgeceae cegtgteace gecatecaca tegaceetge 3300
caccacagg cagaagctgt acacactgca ggacaaggcc caagtggctg acgtggtggt 3360
gagcaggtgg ctgagggtca cagtggccgg aggcgtccac atctccgggc tccacactga 3420
gtcggccccg cggcggcagc aggagcagca ggtgcccatc ctggagaagt tttgcttcac 3480
tececacaeg gaggagggt geetgtetga geacgetgee etegaggagg agetgeaaet 3540
gtgcaagggg ctggtcgagg cactcgagac caaggtgacc cagcaggggc tgaagatggt 3600
ggtgcccgga ctggatgggg cccagatccc cccgggaccc ctcacagcag gaactgcccc 3660
gactattate gactacetae agastteaga teangagan annount guartacetae agastteaga teangagan
ggctgttgtc ggctgcctgc aggcttcagc tcaacgggaa cctgcagctg gagctggcgc 3720
aggtgetgge ccaggagag cccaagetge cagaggacce tetgeteage ggeeteetgg 3780
acteceegge acteaaggee tgeetggaea etgeegtgga gaacatgeee ageetgaaga 3840
tgaaggtggt ggaggtgctg gccggccacg gtcacctgta ttcccgcatc ccaggcctgc 3900
transporter consequent adartises carried to the consequence of the con
tggaggetge ccaggeegag etgeageage acgaegttge ccagggeeag tgggateeeg 4020
cagaccetge ecceagege etgggeageg eggaceteet ggtgtgcaac tgtgetgtgg 4080
ctgccctcgg ggacccgcct cagctctcag caacatggtg gctgccctga nagaaggggg 4140
ctttctgctc ctgcacacac tgctccgggg gcaccccctc ggggacatcg tggccttcct 4200
cacctccact gagccgcagt atggccaggg catcctgagc caggacgcgt gggagagcct 4260
cttctccagg gtgtcgctgc gcctggtggg cctgaagaag tccttctacg gctccacgct 4320
cttcctgtgc cgccggccca ccccgcagga cagccccatc ttcctgccgg tggacgatac 4380
cagetteege tgggtggagt etetgaaggg cateetgget gacgaagaet ettteeegge 4440
ctgtgtggct gaaggccatc aactgttcca cctcgggcgt ggtgggcttg gtgaactgtc 4500
tccgccgaga gcccggcgga acgctccggt gtgtgctgct ctccaacctc agcagcacct 4560

cccacgtcc	c ggaggtgga	c ccgggctcc	g cagaactgc	a gaaggtgtt	g cagggagaco	4620
tggtgatga	a cgtctaccg	c gacggggcc	t ggggggctt	t ccgccactto	ctgctggagg	4680
aggacaagc	c tgaggagcc	g acggcacate	g cctttgtga	g caccctcac	c cggggggacc	4740
tgtccctcc	a tccgctggg	t ctgctcctc	g ctgcgccate	g cccagcccad	ctgccctggc	4800
gcccagctc	t gcacggtct	a ctacgcctc	ctcaacttc	c gcgacatcat	gctggccact	4860
	t cccctgatg					
	t cgggccgaga					
	a cctctgtcct					
	g aggcggcct					
	c gggtgcncc					
	g ccatcgccat					
	a agcgggcgta					
	c gggacacato					
	g tcttgaacto					
	g gtcgcttcct					
	a tcttcctgaa					
	a gtgctgactg					
	ggcccctcaa					
cgctacatg	g cccaagggaa	gcacattggc	aaagtcgtcg	tgcaggtgct	tgcggaggag	5700
ccggaggcag	g tggctgaagg	gggccaaacc	caagctgatg	teggecatet	ccaagacctt	5760
	cacaagagct					
ggcgcagtgg	g ctgatacage	gtggggtgca	gaagctcgtg	ttgacttctc	gctccgggat	5880
	taccaggcca					
	agcaacatca					
	gcccgtgggc					
tggagaacca	gaccccagag	ttcttccagg	acgtctgcaa	gcccaagtac	agcggcaccc	6120
tgaacctgga	cagggtgacc	cgagggcgtg	ccctgagctg	gactactttg	tggtcttctc	6180
ctctgtgagc	tgcgggcgtg	gcaatgcggg	acagagcaac	tacggctttg	ccaatttccg	6240
ccatggagcg	tatctgtgag	aaacgccggc	acgaaggcct	cccaggcctg	gccgtgcagt	6300
ggggcgccat	cggcgacgtg	ggcattttgg	tggagacgat	gagcaccaac	gacacgatcg	6360
tcagtggcac	gctgccccag	cgcatggcgt	cctgcctgga	ggtgctggac	ctcttcctga	6420
accagcccca	catggtcctg	agcagctttg	tgctggctga	gaaggctgcg	gcctataggg	6480
acagggacag	ccagcgggac	ctggtggagg	ccgtggcaca	catcctgggc	atccgcgact	6540
tggctgctgt	caacctggac	agctcactgg	cggacctggg	cctggactcg	ctcatgagcg	6600
tggaggtgcg	ccagacgctg	gagcgtgagc	tcaacctggt	gctgtccgtg	cgcgaggtgc	6660
ggcaactcac	gctccggaaa	ctgcaggagc	tgtcctcaaa	ggcggatgag	gccagcgagc	6720
tgggcatgcc	ccacgcccaa	ggaggatggt	ctggcccagc	agcagactca	gctgaacctg	6780
cgctccctgc	tggtgaaccc	ggagggcccc	accctgatgc	ggctcaactg	ccgtgcagag	6840
ctcggagcgg	cccctgttcc	tggtgcaccc	aattcgaggg	ctccaccacc	gtgttccaca	6900
gcctggcctc	ccggctcagc	atccccacct	atggcctgca	gtgcacccga	gctgcgcccc	6960
ttgacagcat	ccacageetg	gctgcctact	acatcgactg	catcaggcag	gtgcagcccg	7020
agggccccta	ccgcgtggcc	ggctactcct	acggggcctg	cgtggccttt	gaaatgtgct '	708 <b>0</b>
	ggcccagcag					
	ctacgtactg					
	ggctgagacg					
	ggtgctggag					
	cctgatcatc					
cggcccggtc	cttctactac	aagctgcgtg	ccgctgagca	gtacacaccc	aaggccaagt (	7440

```
accatggcaa cgtgatgcta ctgcgcgcca agacgggtgg cgcctacggc gaggacctgg 7500
  gcgcggacta caacctctcc caggtatgcg acgggaaagt atccgtccac gtcatcgagg 7560
  gtgaccaccg cacgctgctg gagggcagcg gcctggagtc catcatcagc atcatccaca 7620
  getecetgge tgagecaege gtgagegtge gggagggeta ggeeegtgee eeegeetgee 7680
  accggaggtc actccaccat ccccacccca tcccacccca cccccgccat gcaacgggat 7740
  tgaagggtcc tgccggtggg accetgtccg gcccagtgcc actgcccccc gaggctagct 7800
  agacgtaggt gttaggcatg tcccacccac ccgccgcctc ccacggcacc tcggggacac 7860
  cagagetgee gaettggaga eteetggtet gtgaagagee ggtggtgeee gtgeeegeag 7920
  gaactggggc tgggcctcgt gcgcccgtgg ggtctgcgct tggtctttct gtgcttggat 7980
  ttgcatattt attgcattgc tggtagagac ccccaggcct gtccaccctg ccaagactcc 8040
  traggrageg tgtgggtccc gcactetgcc cccatttccc cgatgtcccc tgcgggcgcg 8100
  ggcagccacc caagcctgct ggctgcggcc ccctctcggc caggcattgg ctcagcccgc 8160
  tgagtggggg gtcgtgggcc agtccccgag gactgggccc ctgcacaggc acacagggcc 8220
 cggccacacc cagcggcccc ccgcacagcc acccgtgggg tgctgccctt atgcccggcg 8280
 ccgggcacca actccatgtt tggtgtttgt ctgtgtttgt ttttcaagaa atgattcaaa 8340
 ttgctgcttg gattttgaaa tttactgtaa ctgtcagtgt acacgtctgg accccgtttc 8400
 atttttacac caatttggta aaaatgctgc tctcagcctc ccacaattaa accgcatgtg 8460
 atctccaaaa
                                                                    8470
 <210> 11
 <211> 812
 <212> DNA
 <213> Homo sapiens
 <400> 11
 geogeageea atcagegege gtgeoeggge ecetgegtet ettgegteaa gaeggeogtg 60
 ctgagcgaat gcaggcgact tgcgagctgg gagcgattta aaacgctttg gattcccccg 120
 gcctgggtgg ggagagcgag ctgggtgccc cctagattcc ccgccccgc acctcatgag 180
 ccgaccctcg gctccatgga gcccggcaat tatgccacct tggatggagc caaggatatc 240
 gaaggettge tgggageggg aggggggggg aatetggteg eccaeteece tetgaceage 300
 cacccagegg egectaeget gatgeetget gteaactatg ecceettgga tetgeeagge 360
teggeggage gecaaageaa tgccacccat geeetggggt geeecagggg acgteeccag 420
ctcccgtgcc ttatggttac tttggaggcg ggtactactc ctgccgagtg tcccggagct 480
cgctgaaacc ctgtgcccag gcagccaccc tggccgcgta ccccgcggag actcccacgg 540
ccggggaaga gtaccccagc cgccccactg agtttgcctt ctatccggga tatccgggaa 600
cctaccagcc tatggccagt tacctggacg tgtctgtggt gcagactctg ggtgctcctg 660
gagaaccgcg acatgactcc ctgttgcctg tggacagtta ccagtcttgg gctctcgctg 720
gtggctggaa cagccagatg tgttgccagg gagaacagaa cccaccaggt cccttttgga 780
aggcagcatt tgcagactcc agcgggcagc ac
<210> 12
<211> 2385
<212> DNA
<213> Homo sapiens
<400> 12
ataagetggg gtaaagtatt ttegeagttt etgeetttag gattttatta gettetetee 60
cccaggccgc agccaatcag cgcgcgtgcc cgggcccctg cgtctcttgc gtcaagacgg 120
```

```
ccgtgctgag cgaatgcagg cgacttgcga gctgggagcg atttaaaacg ctttggattc 180
 ccccggcctg ggtggggaga gcgagctggg tgcccctag attccccgcc cccgcacctc 240
 atgageegae ceteggetee atggageeeg geaattatge cacettggat ggageeaagg 300
 atatcgaagg cttgctggga gcgggagggg ggcggaatct ggtcgccac tcccctctga 360
 ccagccaccc ageggegeet aegetgatge etgetgteaa etatgeeece ttggatetge 420
 caggetegge ggageegeea aageaatgee acceatgeee tggggtgeee caggggaegt 480
 ecceagetee egtgeettat ggttaetttg gaggegggta etaeteetge egagtgteee 540
 ggageteget gaaaccetgt geccaggeag ceaccetgge egegtacece geggagaete 600
 ccacggccgg ggaagagtac cccagccgcc ccactgagtt tgccttctat ccgggatatc 660
 cgggaaccta ccagcctatg gccagttacc tggacgtgtc tgtggtgcag actctgggtg 720
 ctcctggaga accgcgacat gactccctgt tgcctgtgga cagttaccag tcttgggctc 780
 tegetggtgg etggaacage cagatgtgtt gecagggaga acagaaceca ecaggteeet 840
 tttggaaggc agcatttgca gactccagcg ggcagcaccc tcctgacgcc tgcgcctttc 900
 gtcgcggccg caagaaacgc attccgtaca gcaaggggca gttgcgggag ctggagcggg 960
 agtatgcggc taacaagttc atcaccaagg acaagaggcg caagatctcg gcagccacca 1020
gcctctcgga gcgccagatt accatctggt ttcagaaccg ccgggtcaaa gagaagaagg 1080
 ttctcgccaa ggtgaagaac agcgctaccc cttaagagat ctccttgcct gggtgggagg 1140
agcgaaagtg ggggtgtcct ggggagacca ggaacctgcc aagcccaggc tggggccaag 1200
gactetgetg agaggeeeet agagacaaca ceetteecag geeactgget getggactgt 1260
tcctcaggag cggcctgggt acccagtatg tgcagggaga cggaacccca tgtgacagcc 1320
cactccacca gggttcccaa agaacctggc ccagtcataa tcattcatcc tgacagtggc 1380
aataatcacg ataaccagta ctagctgcca tgatcgttag cctcatattt tctatctaga 1440
gctctgtaga gcactttaga aaccgctttc atgaattgag ctaattatga ataaatttgg 1500
aaggegatee etttgeaggg aagetttete teagaceeee tteeattaca eeteteacee 1560
tggtaacagc aggaagactg aggagagggg aacgggcaga ttcgttgtgt ggctgtgatg 1620
teegtttage attititetea getgaeaget gggtaggtgg acaattgtag aggetgtete 1680
ttcctccctc cttgtccacc ccatagggtg tacccactgg tcttggaagc acccatcctt 1740
aatacgatga tttttctgtc gtgtgaaaat gaagccagca ggctgcccct agtcagtcct 1800
tccttccaga gaaaaagaga tttgagaaag tgcctgggta attcaccatt aatttcctcc 1860
cccaaactct ctgagtcttc ccttaatatt tctggtggtt ctgaccaaag caggtcatgg 1920
tttgttgagc atttgggatc ccagtgaagt agatgtttgt agccttgcat acttagccct 1980
tcccaggcac aaacggagtg gcagagtggt gccaaccctg ttttcccagt ccacgtagac 2040
agattcacgt gcggaattct ggaagctgga gacagacggg ctctttgcag agccgggact 2100
ctgagaggga catgagggcc tetgeetetg tgtteattet etgatgteet gtacetggge 2160
teagtgeeeg gtgggaetea teteetggee gegeageaaa geeagegggt tegtgetggt 2220
cetteetgea cettaggetg ggggtggggg geetgeegge geatteteea egattgageg 2280
cacaggeetg aagtetggae aaceegeaga acegaagete egageagegg gteggtggeg 2340
agtagtgggg tcggtggcga gcagttggtg gtgggccgcg gccgc
                                                                  2385
<210> 13
<211> 221
<212> DNA
<213> Homo sapiens
<400> 13
dsdnrstatc tttctgtgtg gtgcagccct gttggcagtg ggcatctggg tgtcaatcga 60
tggggcatcc tttctgaaga tcttcgggcc actgtcgtcc agtgccatgc agtttgtcaa 120
```

cgtgggctac ttcctcatcg cagccggcgt tgtggtcttt gctcttggtt tcctgggctg 180

```
ctatggtgct aagactgaga gcaagtgtgc cctcgtgacg t
                                                                   221
  <210> 14
  <211> 1533
  <212> DNA
  <213> Homo sapiens
  <400> 14
  gggcacgcag acattctggg aagccacttg ccccacccct gggctgcttc ttcttgagat 60
  caggaggggc gttgcccagg gctggtgttg ccaggtggag gcctgctgag gcagtggttg 120
  tggggatcgg tctccaggca gcaggggca gcagggtcaa ggagaggcta actggccacg 180
 ggtggggcca gcaggcgggc agaaggaggc tttaaagcgc ctaccctgcc tgcaggtgag 240
 cagtggtgtg tgagagccag gccgtccctc tgcctgccca ctcagtggca acacccggga 300
 gctgttttgt cctttgtgga gcctcagcag ttccctgctt tcagaactca ctgccaagag 360
 ccctgaacag gagccaccat ggcagtgctt cagcttcatt aagaccatga tgatcctctt 420
 caatttgctc atctttctgt gtggtgcagc cctgttggca gtgggcatct gggtgtcaat 480
 cgatggggca tcctttctga agatcttcgg gccactgtcg tccagtgcca tgcagtttgt 540
 caacgtgggc tacttcctca tcgcagccgg cgttgtggtc tttgctcttg gtttcctggg 600
 ctgctatggt gctaagactg agagcaagtg tgccctcgtg acgttcttct tcatcctcct 660
 cctcatcttc attgctgagg ttgcagctgc tgtggtcgcc ttggtgtaca ccacaatggc 720
 tgagcacttc ctgacgttgc tggtagtgcc tgccatcaag aaagattatg gttcccagga 780
 agacttcact caagtgtgga acaccaccat gaaagggctc aagtgctgtg gcttcaccaa 840
 ctatacggat tttgaggact caccctactt caaagagaac agtgcctttc ccccattctg 900
 ttgcaatgac aacgtcacca acacagccaa tgaaacctgc accaagcaaa aggctcacga 960
 ccaaaaagta gagggttgct tcaatcagct tttgtatgac atccgaacta atgcagtcac 1020
 cgtgggtggt gtggcagctg gaattggggg cctcgagctg gctgccatga ttgtgtccat 1080
 gtatetgtae tgeaatetae aataagteea ettetgeete tgeeactaet getgeeacat 1140
 caatgtcact tgggccagaa tggacctgcc ctttctgctc cagacttggg gctagatagg 1260
 gaccacteet tttaggegat geetgaettt eetteeattg gtgggtggat gggtgggggg 1320
 cattecagag cetetaaggt agceagttet gttgeecatt ecceeagtet attaaaceet 1380
 tgatatgccc cctaggccta gtggtgatcc cagtgctcta ctgggggatg agagaaaggc 1440
attttatagc ctgggcataa gtgaaatcag cagagcctct gggtggatgt gtagaaggca 1500
cttcaaaatg cataaacctg ttacaatgtt gcc
<210> 15
<211> 472
<212> DNA
<213> Homo sapiens
<400> 15
tcagagaaaa ctcaaacttt attgagagaa ttttcaaatt ttcagtcaca ttttcaatgt 60
gacatcagec atgtgtgtag cttcagettg tettetttt aacttatgge tgeecatete 120
ctgcttcttt agtcttagca tgcttaggat taggtggagt cttctctttt acatcagagc 180
catetecaeg eteaeteega gtetttteea gateeattte etggeaatea eettetaett 240
tacgttette gateggaggt gtteettete tetettgtee aggtteaata teetgattgt 300
cagttggtgg ttcctcttgc tgagattcac cgggagccac gaatgcaacc acatcgggag 360
ceteetgace ateteetett cetetggate ttgateteae tegtgeacte ategetgeaa 420
```

```
ctagaagatc gtgaactgaa gaacttgagt cagcagagag cctggcgaag aa
                                                                    472
 <210> 16
 <211> 478
 <212> DNA
 <213> Homo sapiens
 <400> 16
 cttcattctt cgccaggctc tctgctgact caagttcttc agttcacgat cttctagttg 60
 cagcgatgag tgcacgagtg agatcaagat ccagaggaag aggagatggt caggaggctc 120
 ccgatgtggt tgcattcgtg gctcccggtg aatctcagca agaggaacca ccaactgaca 180
 atcaggatat tgaacctgga caagagagag aaggaacacc tccgatcgaa gaacgtaaag 240
 tagaaggtga ttgccaggaa atggatctgg aaaagactcg gagtgagcgt ggagatggct 300
 ctgatgtaaa agagaagact ccacctaatc ctaagcatgc taagactaaa gaagcaggag 360
 atgggcagcc ataagttaaa aagaagacaa gctgaagcta cacacatggc tgatgtcaca 420
 ttgaaaatgt gactgaaaat ttgaaaattc tctcaataaa gtttgagttt tctctgaa
 <210> 17
 <211> 198
 <212> DNA
 <213> Homo sapiens
<220>
<221> unsure
<222> (191)
<400> 17
cccgctgtac caccccagca tgttctgcgc cggcggaggg caagaccaga aggactcctg 60
caacggtgac tetggggggc ceetgatetg caacgggtac ttgcagggec ttgtgtettt 120
cggaaaagcc ccgtgtggcc aagttggcgt gccaggtgtc tacaccaacc tctgcaaatt 180
cactgagtgg nattaagg
                                                                   198
<210> 18
<211> 465
<212> DNA
<213> Homo sapiens
<400> 18
tggagatgga gtatgtattt attttacaaa aataaatcac catcttcgga ccatttgtag 60
actggaacat ttcgagcaat gagtgcgcca cacggacgag tgccctggtg actccctgat 120
gttcgcgtca cccccagggc caccttggcg cccgcatgag cctcgcttcc cactcccggc 180
ctccaactcc cttccctcgc agccgccatt caccttctgc tgtttatttg tctgcagagc 240
gcctggacac cggaaaaggc gattccctga gcgcctggag ttggagacaa ttcctggttc 300
agaatttaaa catctttcta aggtaagege tgeteeaaaa etettegeeg egtggggaet 360
ttgcaccagg ggcggttggg aaggaagttg gccctccacg ggttcctggg caaccgcggc 420
ctgttgaaaa aaggttctgg gtcaaataat ttaacttcgg aggag
                                                                   465
<210> 19
```

```
WO 00/23111
                                                               PCT/US99/24331
   <211> 204
   <212> DNA
  <213> Homo sapiens
  <400> 19
  ggcgggaaca ggcggcgctg gacctgtacc cctacgacgc cgggacggac agcggcttca 60
  cetteteete ecceaactte gecaceatee egeaggacae ggtgacegag ataaegteet 120
  cctctcccag ccacccggcc aactccttct actacccgcg gctgaaggcc ctgcctccca 180
  tcgccagggt gacactggtg cggc
                                                                      204
  <210> 20
  <211> 294
  <212> DNA
  <213> Homo sapiens
  <220>
  <221> unsure
 <222> (287)
 <400> 20
 gagatttctc ttcaatggct tcctgtgagc tagagtttga aaatatctta aaatcttgag 60
 ctagagatgg aagtagcttg gacgattttc attatcatgt aaatcgggtc actcaagggg 120
 ccaaccacag ctgggagcca ctgctcaggg gaaggttcat atgggacttt ctactgccca 180
 aggttctata caggatataa aggtgcctca cagtatagat ctggtagcaa agtaagaaga 240
 aacaaacact gatctctttc tgccacccct ctgacccttt ggaactnctc tgac
 <210> 21
 <211> 22
 <212> DNA
 <213> Artificial Sequence
 <220>
<223> Description of Artificial Sequence:Synthetic
<400> 21
atcagaacaa agaggctgtg tc
                                                                    22
<210> 22
<211> 21
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Synthetic
<400> 22
atctctaaag ccccaacctt c
                                                                   21
```

WO 00/23111	PCT/US99/24331
<210> 23	
<211> 19	
<212> DNA	•
<213> Artificial Sequence	•
<220>	•
<223> Description of Artificial Sequence:Synthetic	
<400> 23	
tgccgaagag gttcagtgc	19
<210> 24	
<211> 22	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence:Synthetic	
<400> 24	
gccacagtgg tactgtccag at	22
gecacaging tacinical at	22
<210> 25	
<211> 21	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence:Synthetic	
<400> 25	
gctgcaagtt ctccacattg a	21
<210> 26	
<211> 18	
<212> DNA	
<213> Artificial Sequence	
<220>	
223> Description of Artificial Sequence:Synthetic	
versity bescription of Artificial Sequence: Synthetic	
:400> 26	
cageegeagg tgaaacae	18
:210> 27	
2210> 27	
2212> DNA	
213> Artificial Sequence	

<220> <223> Description of Artificial Sequence:Synthetic <400> 27 tggctttgaa ctcagggtca 20 <210> 28 <211> 20 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Synthetic <400> 28 cggatgcacc tcgtagacag 20 <210> 29 <211> 20 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Synthetic <400> 29 cggcaacctg gtagtgagtg 20 <210> 30 <211> 22 <212> DNA <213> Artificial Sequence <223> Description of Artificial Sequence:Synthetic <400> 30 cgcagctcct tgtaaacttc ag 22 <210> 31 <211> 20 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Synthetic

WO 00/23111	PCT/US99/24331
<400> 31	
cgggaaccta ccagcctatg	20
<210> 32	
<211> 20	<b>3</b> .
<212> DNA	
<213> Artificial Sequence	
<220>	•
<223> Description of Artificial Sequence:Synthetic	
<400> 32	
caggcaacag ggagtcatgt	20
<210> 33	
<211> 18	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence:Synthetic	
<400> 33	
tgggcatctg ggtgtcaa	18
<210> 34	
<211> 19	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence:Synthetic	
<400> 34	
cggctgcgat gaggaagta	19
<210> 35	
<211> 22	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence:Synthetic	•
<400> 35	
gcccatctcc tgcttcttta gt	22
<210× 36	

PCT/US99/24331
•
•
21

### INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/24331

A. CLASSIFICATION OF SUBJECT MATTER			
IPC(7) :Please See Extra Sheet.			
US CL :Please See Extra Sheet,			
According to International Patent Classification (IPC) or to bot	h national classification and IPC	<u> </u>	
B. FIELDS SEARCHED		·····	
Minimum documentation searched (classification system follow	ved by classification symbols)		
U.S. : 424/130.1, 141.1, 155.1, 183.1; 435/6, 7.1, 7.23, 7	.9, 91.2; 436/501, 504, 505, 547; 514/44	; 536/23.5	
Documentation searched other than minimum documentation to t	he extent that such documents are include	d in the fields searched	
Electronic data base consulted during the international search (		e, search terms used)	
Medline, Biosis, Embase, Cancerlit, Scisearch, WPIDS, USPA search terms: CSG, cancer specific gene, cancer, diagnosis	ATFULL	, <u>,.</u>	
C. DOCUMENTS CONSIDERED TO BE RELEVANT			
Category* Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.	
Y Database SCISEARCH, Accession Nu Reverse transcriptase-polymerase cha cancer. Urologic Clinics of North Am 2, pages 367-&.	in reaction assays for prostate	1-6	
CHO-CHUNG et al. Antisense Oligonucleotides for the treatment of cancer. Current Opinion in Therapeutic Patents. 1993, Vol. 3, No. 12, pages 1737-1750, see entire document.  BUSSEMAKERS et al. DD3: A new prostate-specific gene, highly overexpressed in prostate cancer. Cancer Research. 01 December 1999, Vol. 59, No. 23, pages 5975-5979.			
Further documents are listed in the continuation of Box (	C. See patent family annex.		
<ul> <li>Special categories of cited documents:</li> <li>'A' document defining the general state of the art which is not considered to be of particular relevance</li> </ul>	"T" later document published after the int date and not in conflict with the app the principle or theory underlying th	lication but cited to understand	
earlier document published on or after the international filing date  "X"  document of particular relevance; the claime considered novel or cannot be considered to in when the document is taken alone cited to establish the publication date of another citation or other			
special reason (as specified)  "Y"  document of particular relevance; the claimed invention cannot  considered to involve an inventive step when the document  document referring to an oral disclosure, use, exhibition or other  "O"  document referring to an oral disclosure, use, exhibition or other  combined with one or more other such documents, such combina			
P* document published prior to the international filing date but later than the priority date claimed	being obvious to a person skilled in  '&'  document member of the same paten		
Date of the actual completion of the international search Date of mailing of the international search report			
10 FEBRUARY 2000 07 MAR 2000			
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	Authorized offices from Sauce GEETHA P. BANSAL	unice to	
Facsimile No. (703) 305-3230	Telephone No. (703) 308-0196		

# INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/24331

	PCT/US99/24331
A. CLASSIFICATION OF SUBJECT MATTER: IPC (7):	-
A61K 39/395, 48/00; C12P 19/34; C12Q 1/68; G01N 33/53, 33/574, 33/546, 33/5	67
A. CLASSIFICATION OF SUBJECT MATTER: US CL:	
424/130.1, 141.1, 155.1, 183.1; 435/6, 7.1, 7.23, 7.9, 91.2; 436/501, 504, 505, 54	7; 514/44; 536/23.5